

OPERATOR'S MANUAL SphygmoCor XCEL System v1



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Copyright

SphygmoCor® XCEL

Pulse Wave Analysis Measurement

Pulse Wave Velocity Measurement

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Disclaimer

This manual has been validated and reviewed for accuracy. The instructions and descriptions it contains are accurate for the AtCor Medical product models at the time of this manual's production. However,

succeeding models and manuals are subject to change without notice. AtCor Medical assumes no liability for damages incurred directly or indirectly from errors, omissions or discrepancies between the product and the manual.

This Manual is produced on the assumption that the operator is an experienced user of Windows 7, 8 or 10 Operating Systems.

If the operator is not familiar with the Windows environment, please refer to the Windows User Manual or Windows Online Help.

Trademarks

"SphygmoCor" is a registered trademark of AtCor Medical Pty Ltd.

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Caution

Federal (USA) law restricts this device to sale by or on the order of a physician

Introduction

The SphygmoCor XCEL Central Blood Pressure Measurement System

The SphygmoCor XCEL System is a non-invasive diagnostic tool for the clinical assessment of central blood pressure. The SphygmoCor XCEL System derives the central aortic pressure waveform from cuff pulsations recorded at the brachial artery. Analysis of the waveform provides key parameters including central systolic pressure, central pulse pressure and indices of arterial stiffness such as augmentation pressure and augmentation index. Increased central systolic pressure and augmentation index have been shown to be markers of cardiovascular risk.

The SphygmoCor XCEL Pulse Wave Velocity Measurement System

The SphygmoCor XCEL System also measures the pulse wave velocity of the arterial pulse waveform travelling through the descending aorta to the femoral artery. The aortic pulse wave velocity is detected from carotid and femoral arterial pulses measured non-invasively and simultaneously. The carotid pulse is measured using the tonometer while the femoral pulse is measured through pulsations in a cuff placed around the thigh. Higher carotid to femoral pulse wave velocity indicates a stiffer aorta, which is an indicator of target organ damage.

Indications for Use

The SphygmoCor[®] XCEL System provides a derived ascending aortic blood pressure waveform and a range of central arterial indices. These measurements are provided non-invasively through the use of a brachial cuff. It is to be used on those patients where information related to ascending aortic blood pressure is desired but the risks of cardiac catheterization procedure or other invasive monitoring may outweigh the benefits.

Additionally, the SphygmoCor[®] XCEL System automatically measures Systolic blood pressure and Diastolic blood pressure.

The SphygmoCor[®] XCEL Pulse Wave Velocity (PWV) option is intended to obtain PWV measurements. The PWV option is used on adult patients only.

Intended Patient Population

The SphygmoCor[®] XCEL System is intended to be used on adult patients only.

Do not use on neonates.

Safety and effectiveness in children under the age of 18 years of age and neonates have not been established.

Intended Environment

The SphygmoCor[®] XCEL System is intended to be used in a Clinical or Research environment.

The SphygmoCor XCEL System is intended to be used temporarily as a cardiovascular diagnostic tool. It is not intended to be used as a continuous monitoring device in hospital or emergency vehicle, and is not applicable to be classified as a defibrillation-proof applied part.

The SphygmoCor XCEL system is not intended for use during patient transport outside of a healthcare facility.

Contraindications

- The SphygmoCor XCEL System should not be used for patients with erratic, accelerated or mechanically controlled irregular heart rhythms, including patients with arrhythmias.
- Do not use the SphygmoCor XCEL System on patients with carotid or aortic valve stenosis.
- The device should not be used on patients with peripheral artery disease PAD or leg artery disease since an applied cuff pressure on the leg of these subjects may be harmful.
- On hypotensive patients, the device should be used with caution.
- The system is not applicable in generalised constriction or localised spasm of muscular conduit arteries such as seen immediately after hypothermic cardiopulmonary bypass surgery or accompanying Raynaud's phenomena or intense cold.
- Any interpretations made from the SphygmoCor XCEL System measurements should be made in conjunction with all other available medical history and diagnostic test information about a patient.
- Do not use the cuff over a wound as this may cause further injury.
- Do not place the cuff on any limb being used for intravenous access or where an arterio-venous (A-V) shunt is present, or any area where circulation is compromised or has potential to be compromised and could result in injury to the patient
- It is advisable for patients with mastectomy not to perform measurement with the cuff on the side of mastectomy as it may cause harmful injury to the patient.
- The user need to check that application of the cuff do not result in prolonged impairment of patient blood circulation

Warnings

- Do not use mobile/cellular phones or other transmitting devices within 10 meters (30 feet) of the SphygmoCor XCEL System.
- Use only with the tonometer, cuffs and air hose supplied by AtCor Medical. Different tonometers, hoses and cuffs have not been validated with SphygmoCor XCEL and measurements with non-validated components may not be accurate,
- Do not use the Tonometer on moist or wet skin.
- Do not put objects on top of the SphygmoCor XCEL device that can obstruct access to the Stop Button.
- Continuous CUFF pressure due to connection tubing kinking, result in continuous disruption blood flow may cause harmful injury to the patient.
- Too frequent cuff PWA measurements can cause injury to the patient due to blood flow interference
- Do not modify this equipment without authorisation of the manufacturer.
- Temperature of tonometer and cuff may rise above 41 degree Celsius if operating device at maximum operating ambient temperature.
- To disconnect from the supply mains, the power adaptor needs to be unplugged from the mains socket. Ensure the device is positioned such that the power adaptor plug (supply mains side) is not obstructed during use

• Note: additional warnings printed on the SphygmoCor XCEL device and at the end of the Manual.

SphygmoCor XCEL PWA Quick Start Guide

This Quick Start Guide is based on default software options for SphygmoCor XCEL. Changing from default settings may result in different menu options.

1. Software Installation

Note: Make sure that the SphygmoCor XCEL device is NOT connected during the software installation.

Refer to section Software Download

The SphygmoCor XCEL software installation media can be downloaded from the AtCor Medical website.

- 1. Open atcormedical.com in your web browser.
- 2. Navigate to the **Customer Support** section.
- 3. Create an account.
- 4. You will now have access to the XCEL Software, User Manual and Inservice videos.
- 5. To download the required files, right click on each file and select 'Save link as'. Save the files to your local drive.
- 6. To open the XCEL Software zip file, right click on the zip file and extract all files to your PC.

Software Installation on *Page 57* for instructions to install the XCEL software.

2. Hardware Installation

a) Connect the power supply to the SphygmoCor XCEL device and plug in the power supply into a mains socket.

Warning: Do not connect any power supply to the SphygmoCor XCEL device other than the one provided by AtCor.

- b) Connect the SphygmoCor XCEL device to your PC using the USB cable.
- c) Connect the brachial cuff to the SphygmoCor XCEL device.
- d) Switch on the SphygmoCor XCEL device.
- e) Fit the cuff to the patient
- f) Start the SphygmoCor XCEL software by double-clicking on the SphygmoCor XCEL software icon
 f) on the desktop.

Note: If the following error (Code=3251) appears after starting the SphygmoCor XCEL software, refer to *Installing Recommended Fonts on Different Versions of Windows* section in the *Appendix*.



3. Creating a Patient Record

a) Click **New** to add a new patient record, or go to **b** if it is the first patient record.

					Hide
Р	atient ID:			Date	of Birth*: Month Day Year
Fir	st Name:				Gender*:Select
Las	t Name*: Subject				
New	Search	Edit	Delete	Save	Cancel

b) Enter all mandatory data, indicated by the * symbol next to the field.

Note: Before creating a new patient entry, check whether the patient already exists in the database. Separate patient entries cannot be merged.

c) Click Save.

4. Recording a PWA Measurement

Note: Minimise the patient's limb movement while capturing data. Movement may prevent the capture from being completed or may reduce the quality of the measurement.

a) Select PWA (this is only necessary if you also have a license for PWV).

2		Syster	n Database	Help			
	Mode	PWA	•			Setu	• Capture F
		PWV		Patients 5			
	Patier	nt ID	La	st Name	First Name	Date Of Birth	Gender
	1		Smith		John	1/1/1954	Male
	2		Doe		Jane	4/7/1970	Female
	3	Roe		Jane	10/16/1966	Female	
	4		Smith		Johnnie	7/8/1998	Male
	5		Doe		Janie	12/30/1997	Female



b) Select the patient (search for the record if necessary)

c) Click Start.

The cuff is inflated to measure patient's brachial systolic pressure (SYS), diastolic pressure (DIA) and pulse pressure (PP) which will be displayed. During a BP assessment, the pressure is shown both with a bar, and a numerical value on the right side of the screen.



Note: If the multiple brachial BP assessments option is selected in the Brachial BP setting, the last displayed brachial SYS, DIA and PP will be the average values of the successfully completed assessments.

Five seconds after the cuff deflation, the PWA measurement starts automatically.



Note: When using the multiple brachial BP assessments option, there may be occasions where not all Brachial BP assessments were performed successfully. On these occasions, the user can either start the PWA measurement manually by clicking the Capture button or can start a new multiple Brachial BP assessment by clicking the Start button.



The cuff is inflated and the capture screen is displayed, showing the captured cuff waveform.



The cuff can be deflated at any time by clicking the **Cancel** button on the screen, manually pressing the **Stop** button on top of the SphygmoCor XCEL device or turning off the SphygmoCor XCEL device.

Note: SphygmoCor XCEL inflates the cuff before each reading, then deflates the cuff.

d) After 5 seconds the recorded cuff waveform will be automatically captured and a report screen will be displayed showing the central aortic pressure waveform and parameters.

Refer to the *Using the Guidance Bars* section on *page 24* for explanation of the bars in the capture screen.



Note: If the report screen does not display after 20 seconds of recorded brachial cuff pulses, click the **Capture** button to produce a report.

Refer to *Reviewing the PWA Report* section on *page* 25 for an explanation of the report screen displayed parameters.

- e) Click **Print** for a single-page summary report or click **Print Patient Report** for a multi-page patient report. The patient report may not be available in some regions.
- f) Click Repeat to repeat the measurement on the same subject.
- g) Click **Delete** to delete the report.

SphygmoCor XCEL PWV Quick Start Guide

This Quick Start Guide is based on default software options for SphygmoCor XCEL. Changing from default settings may result in different menu options.

1. Software Installation

Note: Make sure that the SphygmoCor XCEL device is NOT connected during the software installation.

Refer to Software Download

The SphygmoCor XCEL software installation media can be downloaded from the AtCor Medical website.

- 7. Open atcormedical.com in your web browser.
- 8. Navigate to the **Customer Support** section.
- 9. Create an account.
- 10. You will now have access to the XCEL Software, User Manual and Inservice videos.
- 11. To download the required files, right click on each file and select 'Save link as'. Save the files to your local drive.
- 12. To open the XCEL Software zip file, right click on the zip file and extract all files to your PC.

Software Installation section on *Page 57* for instructions to install the XCEL software.

2. Hardware Setup

a) Connect the power supply to the SphygmoCor XCEL device and plug in the power supply into a mains socket.

Warning: Do not connect any power supply to the SphygmoCor XCEL device other than the one provided by AtCor.

- b) Connect the SphygmoCor XCEL device to your PC using the USB cable.
- c) Switch on the SphygmoCor XCEL device.
- d) Fit the cuff on the patient's upper thigh.
- e) Start the SphygmoCor XCEL software.
- 3. Creating a Patient Record
- a) Click **New**.

			Hide
Patient ID:			Date of Birth*: Month Day Year
First Name:			Gender*:Select
Last Name*: <mark>Subj</mark>	ect		
New Search	Edit	Delete	Save Cancel

b) Enter all mandatory data, indicated by the * symbol next to the field.

Note: Before creating a new patient entry, check whether the patient already exists in the database. Separate patient entries cannot be merged.

- c) Click Save.
- 4. Recording a PWV Measurement

Note: Minimise the patient's limb movement while capturing data. Movement may prevent the capture from being completed or reduce quality of the measurement.

a) Go to the setup screen.

b) Select Mode - PWV.

t,	Syster	n Database	Help					
Mode	PWV					Setup	Capture	
	PWV PWA							
Patie	ent ID	Las	t Name	First Name	Date	Of Birth	Gender	
1	1 Smith		John	1/1/19	54	Male		
2	2 Doe		Jane	4/7/19	70	Female		
3		Roe		Jane	10/16/	1966	Female	
1		Smith		lohnnie	7/8/10	98	Male	

- c) Select the patient's record.
- d) Enter the distance in millimetres from the patient's carotid artery to their sternal notch (*Carotid to Sternal Notch*) and also the distance from their sternal notch to the top edge of the femoral cuff (*Sternal Notch to Cuff*).



Note: The femoral to cuff distance value is set to 200mm by default. To improve the accuracy of the PWV reading, measure the distance from the top edge of the cuff to the femoral artery (near the groin) and use this value instead of the default.

Refer to

PWV Settings section on page 77 for details on distance measurements.

- e) Click Capture.
- f) The cuff will be inflated to capture femoral waveform, and the carotid signal should be measured using the tonometer. When both signals are of "good" quality, both signals turn green and a PWV report is produced.



The cuff can be deflated at any time by clicking the **Cancel** button on the screen, manually depressing the **Stop** button on top of the SphygmoCor XCEL device or by turning off the SphygmoCor XCEL device.

Note: SphygmoCor XCEL inflates the cuff before each reading, then deflates the cuff.

g) After 10 seconds of simultaneous valid carotid tonometer signal and femoral cuff signal, a report screen will appear showing both signals and the calculated pulse wave velocity (PWV) and pulse transit time.

Refer to *Reviewing the PWV Report* section on *page 49* for an explanation of the parameters displayed in the report screen.

Note: If the report screen does not appear for over 20 seconds of continuous simultaneous recording of carotid and femoral pulses, click the **Capture** button to produce a report.



- h) Click **Print** for a single-page report or **Print Patient Report** for a multi-page patient report. The patient report may not be available in some regions.
- i) Click Repeat to repeat the measurement on the same subject.
- j) Click **Delete** to delete the report.

Recording a PWA Measurement

Components and Accessories for PWA



Brachial cuff – a pneumatic cuff with a tube connection to the SphygmoCor XCEL device Part numbers: 1-00889 - Cuff PWA Adult Large 31-40cm 1-00890 - Cuff PWA Adult 23-33cm 1-00891 - Cuff PWA Adult Extra Large 38-50cm
Cuff Hose- 1.5m cuff hose with connectors at both ends. Part number: 1-00865 - Cuff Hose 1.5m
Power supply – an AC power adaptor for the SphygmoCor XCEL device. Always use the power supply provided by AtCor Medical Part number: 1-011910877 - Medical Power Adaptor (Mean Well – PN GSM36B15-R7B)
USB cable (2m) - connects the SphygmoCor XCEL device to a PC or laptop. Part number: 1-00858 - USB Cable 2m

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Tray (optional) – provides storage of cuffs, as well as a mounting platform for the SphygmoCor XCEL device (The storage tray is an optional component supplied by AtCor Medical).

Part number: 1-00896 - EM4 Storage Tray

Connect Components

Warning: Do not connect any power supply to the SphygmoCor XCEL device other than the power supply provided by AtCor Medical.

- 1. Connect the power supply to the SphygmoCor XCEL device and plug the power supply into an AC power outlet.
- 2. Connect the SphygmoCor XCEL device to the PC using the USB cable.
- 3. Connect the brachial cuff to the SphygmoCor XCEL device.
- 4. Power on the SphygmoCor XCEL device.

Note: Avoid compression or restriction of the connection tubing.



Start a Measurement

Note: The patient's limb movement should be minimised while capturing data as movement may prevent the capture from being completed or may reduce the quality of the measurement.

3		Syster	n Database	Help			
	Mode	PWA	•			Setu	p Capture F
		PWV PWA		Patients 5			
	Patie	nt ID	Last	Name	First Name	Date Of Birth	Gender
	1		Smith		John	1/1/1954	Male
	2		Doe		Jane	4/7/1970	Female
	3		Roe		Jane	10/16/1966	Female
	4		Smith		Johnnie	7/8/1998	Male
	5		Doe		Janie	12/30/1997	Female

1. Select PWA (this is only necessary if you also have a license for PWV).

 Select the patient (by searching Patient ID, Name or Date of Birth, if necessary). If the patient does not exist in the database, create a new patient record by clicking New and entering the patient details. Mandatory fields are indicated by *.

	Hide	
Patient ID:	Date of Birth*: Month Day Year	
First Name:	Gender*:Select	
Last Name*:	Subject	
New Searc	Edit Delete Save Cancel	

3. To determine the patient's SphygmoCor reference age, an estimate of the arterial age of the patient, enter the height of the patient in the height entry box.





4. Click Start.

The cuff is inflated on the right of the screen, to measure brachial systolic and diastolic pressure. During a BP assessment, the pressure is shown both with a bar, and a numerical value on the right side of the screen.



Brachial Systolic Pressure (SYS), Brachial Pulse Pressure (PP) and Brachial Diastolic Pressure (DIA) are displayed. If multiple Brachial blood pressure measurements are selected to calibrate the waveform, the Brachial BP measurement will be repeated and average values for brachial SYS, PP and DIA will be displayed.





After determining the Brachial Systolic and Diastolic pressure, the cuff will deflate. Five seconds later, the cuff will inflate again and automatically capture the PWA waveform.

Note: During PWA waveform capture, the cuff can be deflated at any time by clicking the **Cancel** button on the screen, manually depressing the **Stop** button on top of the SphygmoCor XCEL device or turning off the SphygmoCor XCEL device.

The Capture screen displays the brachial waveform being collected from the cuff. The colour of the waveform along with vertical bar on the side becomes green when the waveform is consistent for capture. When the waveform colour and the side bar become green, the waveform will be automatically captured and the software will advance to the report screen. The waveform may also be captured manually by clicking the capture button at the bottom of the screen.





5. If the Auto Capture feature is disabled, waveforms are captured by clicking the **Capture** button when it is enabled. The **Capture** button is enabled when the software detects a sufficient number of pulses to produce a valid report.

Note: For best results, click the **Capture** button when the waveform colour and corresponding vertical bar are green, as shown in the following screen shot.



A report screen is then displayed, showing Average Aortic Pressure Pulse, aortic pressure waveform parameters and the Aortic Clinical Parameters plotted in reference to the normal range based on the patient's age and gender.



Refer to *Reviewing the PWA Report* for a detailed explanation of this report and the Aortic pressure and arterial stiffness parameters.

- 6. Click **Print** for a single-page report or click **Print Patient Report** for a multi-page patient report. The patient report may not be available in some regions.
- 7. Click **Repeat** to repeat the measurement on the same subject.
- 8. Click **Delete** to delete the report.

Automatic versus Manual Waveform Capture

Waveform capture can be performed either automatically or manually.

- If Auto Capture is enabled (default setting), the software will automatically capture the waveform when all of the quality parameters are met. Alternatively, the keyboard space bar can be pressed at any time to override the Auto Capture feature and capture the waveform.
- If Auto Capture is disabled, data capture can be completed by manually pressing the spacebar on the computer keyboard when the **Capture** button is enabled, indicating enough pulses have been recorded to produce a valid report.

Using the Guidance Bars

When preparing to capture waveforms, the SphygmoCor XCEL System provides feedback that will help determine if the waveforms from the brachial cuff are consistent enough to provide an accurate central blood pressure (CBP) measurement.

There are multiple visual indications that provide feedback, as shown in the following diagram.

All of these visual signals are colour-coded, with some providing feedback based on each pulse, and some on the entire recorded waveform for the duration of the capture time.

If Auto Capture is enabled, the data is captured automatically when the waveform colour changes to green, indicating an acceptable quality waveform has been captured for the duration of the capture time. A report of calculated CBP and arterial stiffness indices is produced. The waveform can also be captured manually by clicking the **Capture** button, regardless of the waveform colour.





1 Waveform – the waveform changes colour according to quality of the overall recording during capture time. The waveform colour is initially white, changes to green if the quality of the overall recording during the capture time is acceptable and the recording has at least the minimum number of quality pulses. The minimum number of pulses recorded can vary according to the data capture period selected in the PWA settings.

2Top Guidance Bar – a Guidance Bar is displayed at the top of the waveform and indicates reliability of the pulse peaks. It updates on a beat-by-beat basis. The colour changes from red to green as the waveform peaks become more consistent.

3 Bottom Guidance Bar – a Guidance Bar is displayed at the bottom of the waveform and indicates reliability of the pulse troughs. The colour changes from red to green as the waveform troughs become more consistent.

4 Bar graph – this graph indicates signal amplitude, measured on a beat-by-beat basis. The amplitude must be above the threshold mark **5** to ensure a reliable signal.

The waveform colour is synchronised with the bar graph colour:

- when the waveform is green, the bar graph is green
- when the waveform is white, the bar graph is red

Note: Guidance Bars and Auto Capture are part of the Capture Guide feature in the software. Both features are enabled as the default setting. These features can be enabled or disabled from the System menu (refer to

Page 25

PWV Settings, PWA Settings, and Brachial BP Settings).

Reviewing the PWA Report

System Database Help		- E
Mode PWA 💌	Setup Capture Report	SphygmoCor 5
Patient Details John Smith Patient ID: 1 Gender: Male	Aortic Clinical Parameters 112	Average Aortic Pressure Pulse 이 편 같
Date Of Birth: 1/1/1954 Age: 62 Measurement Details Report Interpretation	SP Avrid mmHg 105 128 SP: 112	mmHg
Brachial SYS/DIA: 120 / 80 mmнg QC: ✓ ⇒ Height: 178 cm	PP DP: 80 mmHg 29 47 PP: 32	mmHg 110
Notes:	AP MAP: 93	mmHg
	Alx	bpm av Reflect
Edit	Alx75	70- 200 400 600 800 ms
Assessments: 6/9/2016 10:48 AM 6/9/2016 10:48 AM	SphygmoCor Reference	e Age > 59
6/9/2016 10:49 AM	Delete Repeat	Print Patient Report Print

1 Patient Details – Patient's personal data including Patient ID, Gender, Date of Birth and Age.

Measurement Details –

- a) Brachial SYS/DIA: Measured brachial systolic (SYS) and diastolic (DIA) blood pressure.
- b) Height: Patient's height.
- c) Notes: The notes entered by the operator for the measurement.
- d) Quality Control (QC) Indicator: An indicator of the quality of the recorded signal. The QC indicator will appear as a green tick if the recorded signal is a "good" quality signal and as a red cross if the recorded signal is "bad" quality due to noise or patient movement. If the QC indicator displays a red cross, the assessment should be repeated. Click the arrow button enext to the QC indicator to show the Quality Control report. Refer to Quality Control Report (QC) section for more information.

B Report Interpretation – Interpretation for a report entered by the operator.

Assessments – Chronologically ordered list of measurements where each measurement is stamped with the date and time of the measurement. If the patient has more than one assessment, clicking on any assessment allows viewing of the corresponding report. A different assessment can be selected by pressing the arrow key or the tab key on the keyboard.

5 Report Types – Choose a tab for the report type (Clinical, Detailed, Reference Range, and Wave Reflection) from the right side of the screen. Refer to *PWA Report* Types section for more information.

Deleting Reports

Each measurement is automatically saved in the database. Reports can be deleted, if desired, as follows:

1. Click on the assessment to be deleted. Click **Delete** and the following message will be displayed.





2. Click Yes to delete the assessment.

Note: Reports can be deleted at any time, not only at the time of creation.

Repeat the measurement

Click the **Repeat** button, which will only be enabled at the last assessment. This will perform another measurement on the same patient and create another assessment for the patient.

Print the report

The **Print** buttons provide a choice of printed reports.

Print Patient Report Print

Print – a single-page PWA report of the type selected.

Print Patient Report – a multi-page report that provides an explanation of the different PWA parameters. This report may not be available in some regions.

Quality Control Report (QC)

A QC report of the captured waveform is shown by clicking an ext to the **QC indicator** in the **Report** page. The QC report includes the following information:



1 Captured peripheral (brachial) waveform chart.

2 Derived aortic pressure waveform chart.

3 Overlaid peripheral (brachial) pulses.

Quality Control – Quality control indices quantifying the quality of the captured peripheral (brachial) waveforms listed as following:

- Number of Pulses: The number of pulses considered in the calculation.
- Pulse Height: The average pulse height of all the individual pulses in digital units.
- **Pulse Height Variation:** The average pulse height variation (in %) of all individual pulses from the average pulse.
- **Diastolic Variation:** The average diastolic or baseline variation (in %) of all individual pulses from the average pulse.
- Shape Deviation: The average waveform shape variation (in %) of individual pulses from the average pulse.
- **Pulse Length Variation:** The average pulse length variation (in %) of individual pulses from the average pulse.
- Overall Quality: A parameter quantifying the captured peripheral (brachial) waveform quality based on the Quality Control indices. For an Overall Quality value above or equal to 75%, the QC will be a green tick
 and for an Overall Quality value below 75%, the QC will be a red cross X.

PWA Report Types

Clinical

To view the Clinical report, click the **Clinical** tab on the right side of the Report page. The Clinical report contains the following information:





OAverage Central Aortic Pulse – A graphical representation of the derived average aortic pressure pulse. The shape of the aortic pressure pulse is a result of the ventricular ejection and physical properties of the arterial system. The waveform shape changes with changes in arterial stiffness.

Aortic Parameters – A summary representation of the derived average central pressure waveform parameters: SP (Aortic Systolic pressure) is the highest pressure value recorded in a pulse, DP (Aortic Diastolic pressure) is the lowest pressure value recorded in a pulse, PP (Aortic Pulse Pressure) is the height of the aortic pressure pulse, MAP (Mean Arterial Pressure) is the average pressure in a pulse and HR (Heart Rate) is the number of heart beats per minutes.

3 Aortic Clinical Parameters – A bar graph representing the aortic clinical parameters in reference to a healthy population range based on the patient's age and gender:

- **SP:** Aortic Systolic Pressure (mmHg) is the maximum aortic pressure during cardiac ejection. A high aortic SP indicates high cardiovascular load. High arterial stiffness increases the reflected pressure wave in the arterial system, which augments or increases Aortic Systolic Pressure.
- **PP:** Aortic Pulse Pressure (mmHg) is the height of the aortic pressure pulse. PP can also be described as the difference between the maximum and minimum of the aortic pressure pulse, or the Aortic Systolic pressure minus the Aortic Diastolic pressure. High aortic PP has been shown to predict cardiovascular events¹.

¹Roman MJ, Devereux RB, Kizer JR, *et al.* High central pulse pressure is independently associated with adverse outcome. The Strong Heart Study. J Am CollCardiol 2009;54:1730-4.



- **AP:** Aortic Augmented Pressure (mmHg) is the difference between the two aortic pressure peaks during ejection (systole). AP is a measure of wave reflected back from lower body. The value of AP is affected by both the magnitude and speed of the reflected wave, which is an indicator of arterial stiffness. With aging, the arteries become stiffer, consequently, wave reflection increases, leading to increased Aortic Augmented Pressure and resulting in increased risk of cardiovascular disease or organ damage².
- **Alx:** Augmentation Index (%) is the ratio of AP to PP. Studies have shown that patients with diabetes tend to have high Alx, indicating stiffer arteries and higher risk of organ damage³.
- Alx 75: The Alx is normalised for a heart rate of 75 beats per minute. Alx75 enables comparison of reports from patients with different heart rates, and enables comparison of arterial stiffness between visits of the same patient.

SphygmoCor Reference Age – Estimated arterial age based on the average aortic pressure parameters of a healthy population. The patient's height must be entered on the set up screen, or the message "SphygmoCor XCEL reference age not calculated" will appear.

5 Printing PWA Clinical Report – Click Print for a single-page PWA Clinical Report.

² Chirinos et al Aortic Pressure Augmentation Predicts Adverse Cardiovascular Events in Patients With Established Coronary Artery Disease, Hypertension. 2005;45:980-985

³ London et al Arterial Wave Reflections and Survival in End-Stage Renal Failure, Hypertension. 2001;38:434-438.

Printed PWA Clinical Report Example



Detailed

To view the Detailed report, click the **Detailed** tab on the right side of the Report page. The Detailed report contains the following information:



Average Peripheral Pulse-Brachial – Graphical representations of the average of all brachial pressure pulses in the captured brachial waveform. A summary report of average peripheral pulse parameters is represented on the right side of the chart and includes:

- SYS (Peripheral Systolic Pressure in mmHg): Measured brachial systolic pressure or the highest pressure value in the average peripheral pulse.
- DIA (Peripheral Diastolic Pressure in mmHg): Measured brachial diastolic pressure or the lowest pressure value in the average peripheral pulse.
- PPP (Peripheral Pulse Pressure in mmHg): The height of the average peripheral pulse.
- MAP (Mean Arterial Pressure in mmHg): The mean pressure of the average peripheral pulse.

2 Average Aortic Pressure Pulse – Graphical representations of the average of all aortic pressure pulses in the derived aortic pressure waveform. A summary report of the average aortic pressure pulse parameters is represented on the left side of the chart and includes:

- SP (Aortic Systolic Pressure in mmHg): The highest pressure value in the average aortic pulse.
- DP (Aortic Diastolic Pressure in mmHg): The lowest pressure value in the average aortic pulse.
- PP (Aortic Pulse Pressure in mmHg): The height of the average aortic pulse.



• HR (Average Heart Rate in bpm): The average number of heart beats per minute.

3 PP Amplification (PPP/PP) % – The amplification of the peripheral pulse pressure (PPP) with respect to the aortic pulse pressure (PP).

4 Central Haemodynamic Parameters – Central haemodynamic parameters calculated from Average Aortic Pressure Pulse.



- Heart Rate, Period (HR, T_F) Heart Rate in beats per minutes (bpm) is the patient's average heart rate over the capture time. Period in millisecond (ms) is the length of an averaged derived central pulse.
- Ejection Duration (ED) Ejection Duration is the period of time in milliseconds and in percentage to the full cardiac period from the start of the pulse for which the aortic valve is open (T₀) to the closure of the aortic valve (ED), End of Systole.
- Aortic T2 (in millisecond) Aortic T2 is the duration from the start of the pulse to the 2nd systolic peak (P2).
- **P1 Height (P1-DP) (P1H) P1** Height in mmHg is the Primary wave Pressure at T1 which is the difference between the minimum pressure and the pressure at the first systolic peak (P1).
- Aortic Augmentation (AP) The augmented pressure (AP) in mmHg is the difference between the first systolic peak pressure (P1) and second systolic peak pressure (P2).
- Aortic Alx (AP/PP, P2/P1) Aortic Augmentation Index calculated using two different ratios (in %):
 - Alx (AP/PP) The ratio of the augmented pressure (AP) to the pulse pressure (PP).
 - Aix (P2/P1) The ratio of the reflected peak (second systolic peak pressure P2) to the primary peak (first systolic peak pressure P1)
- Aortic Alx (AP/PP) @HR75 in % Aortic Augmentation Index (Alx (AP/PP)) adjusted to a standard heart rate of 75 beats per minute.

- Buckberg SEVR (SEVR %) Buckberg SEVR ratio also known as the subendocardial viability ratio, is the ratio of the Diastolic Area to the Systolic Area of the pressure pulse reflecting cardiac blood supply to organ's demand ratio.
- **PTI (Systole, Diastole) in mmHg.s/min Pressure** Time Index (PTI) is the area under the pressure pulse. PTI (Systole) is area under the pressure pulse during systole per minute. This area is related to the work of the heart and oxygen consumption. PTI (Diastole) is the area under the pressure pulse during diastole per minute. PTI (Diastole) is associated with the pressure and time for coronary perfusion, and so is related to the energy supply of the heart.
- End Systolic Pressure (ESP) in mmHg The pressure at the end of systole.
- MAP (Systole, Diastole) in mmHg- Mean Arterial Pressure during systole is the mean pressure between T₀ to ED and mean arterial pressure during diastole is the mean pressure between the ED and the end of the averaged pulse (T_F).

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5 Printing PWA Detailed Report – Click Print for a single-page PWA Detailed Report.

Printed PWA Detailed Report Example



Reference Range

To view the Reference Range report, click the **Reference Range** tab on the right side of the Report page. The Reference Range report contains the following information:



The population reference data⁴ graphs associated with each parameter are represented indicating the patient's position relative to the reference population data. On each graph, the green line indicates the reference population mean and the red lines indicate the 90% confidence interval on either side of that mean. The reference range for the Central Systolic Pressure (SP), Central Augmented Pressure (AP), Central Pulse Pressure (PP) and Central Augmentation Index (AIx) are plotted by age (years). The Ejection Duration (ED %) and SEVR (%) are plotted by Heart Rate (bpm).

1 Central Systolic Pressure (SP) – Central Systolic Pressure (SP) is the highest blood pressure measured during an average heartbeat at the heart (not at the arm). This graph shows the patient's central systolic pressure value by patient's age relative to the reference population central systolic pressure from 20-90 years old.

 ⁴ London et al Arterial Wave Reflections and Survival in End-Stage Renal Failure, Hypertension. 2001;38:434-438.
 The Anglo-Cardiff Collaborative Trial (ACCT), Am Coll Cardiol 2005;46:1753–60


2 Central Augmented Pressure (AP) – Central Augmented Pressure (AP) is the additional pressure measured at the heart due to the stiffness of the arteries. The lower AP is indicator of less stiff arteries. This graph shows the patient's central augmented pressure value by patient's age relative to the reference population central augmented pressure from 20-90 years old.

3 Central Pulse Pressure (PP) – Central Pulse Pressure (PP) is the difference between central systolic pressu re (SP) and central diastolic pressure. It represents the change in pressure in central aortic artery. This graph shows the central pulse pressure value by patient's age relative to the reference population central pulse pressure from 20-90 years old.

Central Augmentation Index (AIx) – Central Augmentation Index (AIx) is the amount of augmented pressure (AP) in reference to central pulse pressure (PP). Central Augmentation Index is another measure of arterial stiffness. The lower AIx is an indicator of less stiff arteries. This graph shows the central augmentation index value by patient's age relative to the reference population central augmentation index from 20-90 years old.

5 Ejection Duration (ED %) – The Ejection Duration (ED %) is the duration of the ventricular ejection. It is the period of time from the start of the pulse for which the aortic valve is open to the closure of aortic valve (incisura), end of systole. This graph shows the ejection duration value by patient's heart rate relative to the reference population ejection duration with heart the rate range of 40-100 beats/min.

6 SEVR (%) – The Sub-endocardial Viability Ratio (SEVR %) is the ratio of cardiac blood supply to organ's demand. This graph shows the SEVR value by the patient's heart rate relative to the reference population SEVR with the heart rate range of 40-100 beats/min.

Printing PWA Reference Range Report – Click **Print** for a single-page Reference Range Report.



Printed PWA Reference Range Report Example



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Wave Reflection

To view the Wave Reflection report, click the **Wave Reflection** tab on the right side of the Report page. The Wave Reflection report contains a graphical representation of the aortic pulse including forward and reflected pulses and a flow wave as well as wave reflection parameters.



Aortic Pulses and Flow Wave – The central aortic pressure waveform is shown in blue. This waveform is the sum of two pressure waveforms: the pulse wave generated by cardiac ejection is called Forward Pulse shown in green and the pulse wave reflected from the arteries back to the heart is called Reflected Pulse shown in orange. Aortic flow wave resulting from the separation of forward and reflected pulses is also shown in black.

2 Wave Reflection Parameters – The parameters of the reflected waveform.

- Forward Pulse Height (Pf): The height of forward pulse wave.
- Reflected Pulse Height (Pb): The height of reflected pulse wave.
- **Reflection Magnitude:** The ratio of Reflected Pulse Height to Forward Pulse Height in percentage.

B Printing PWA Wave Reflection Report – Click Print for a single-page PWA Wave Reflection Report.



Printed PWA Wave Reflection Report Example





Viewing PWA Reports

Measurements from previous sessions can be displayed at any time.

- 1. Select the **Report** tab.
- 2. Select the session from the Assessments list.
- 3. Select one of the four report types (Clinical, Detailed, Reference Range, Wave Reflection) from the right side of the screen.

Analysis of Multiple PWA Reports for the Same Patient

The Analysis feature can be used to compare multiple measurements for a selected patient. At least 2 assessments are required to use this feature, and a maximum of 30 measurements can be compared.

- 1. Select PWA.
- 2. Click Report.
- 3. Select the first assessment for comparison (click the corresponding date under the Assessment heading).
- 4. Press **Ctrl** and select another assessment. Check the Analysis screen to make sure the assessments included for analysis are the same as your desired assessments. The number of blue dots on Central SP graph shows the number of assessments included in the analysis, with the the date and time of assessment is noted on the x axis.

Note: To select a new set of assessments for analysis, click once on any of the assessment in the Assessment section and follow the same procedure from step 3

5. The selected assessments are plotted.



6. (Optional) Click **Print** to print a copy of the Analysis report.

7. Select the **Setup** tab to exit the assessment screen or click on any assessment to return to the report screen.

Techniques for Obtaining Quality PWA Measurements

Control Conditions

When performing a SphygmoCor XCEL measurement, it is recommended that the patient:

- Abstain from alcohol for at least 6 hours prior to the measurement.
- Abstain from tobacco and caffeine for at least 4 hours prior to the measurement.
- Fast for at least 6 hours prior to the measurement, however, if this cannot be achieved, a light meal before the measurement is allowed.
- Rest (seated or supine position) comfortably, for at least 5 minutes prior to the measurement.
- Has their back and arm supported during the measurement.
- Be as relaxed as possible and not talk during the measurement
- When possible, subsequent measurements should be obtained at the same time of the day so as to account for the diurnal blood pressure effect.

Note: Blood pressure can be affected by the measurement site, the position of the patient (standing, sitting, lying down), exercise, or the patient's physiological condition.

Cuff Placement

1. Select a cuff that is an appropriate size for the patient's limb.

Each cuff has a size range marked on the inside. Brachial cuffs also have an index mark on the end. When wrapped around the patient's arm the index mark must fall within the marked range. If it is outside this range, select a different cuff.

2. Place the cuff on the upper arm, centred on the brachial artery. Palpate the brachial artery if necessary in order to locate accurately the artery.

The cuff should be placed on bare arm. Remove tight, bulky clothing from the upper arm to prevent constriction



3. Hold the cuff open with the tube pointing downward. Slide the cuff onto the upper arm.



4. Position the cuff with the artery marker over the brachial artery and aligned with the patient's small finger. The cuff should be as high as possible on the arm.



- 5. Fasten the cuff so it is snug, but not too tight. Two fingers should fit between the patient's arm and the cuff.
- 6. Position the arm to ensure that the cuff (i.e., middle of the cuff) and the heart are at the same level.

Note: Avoid compression or restriction of the connection tubing

Recording a PWV Measurement

Components and Accessories for PWV



Femoral cuff (light blue) – a large, thin pneumatic cuff with a connector for the SphygmoCor XCEL device Part number: 1-00897 - PWV Cuff Adult 1-00898 - PWV Cuff Large Adult
Cuff Hose 2m cuff hose with connectors at both ends Part number: 1-00864 - Cuff Hose 2m
Power supply – an AC power adaptor for the SphygmoCor XCEL device. Always use the power supply provided by AtCor Medical with your SphygmoCor XCEL device. Part number: 1-01191 - Medical Power Adaptor (Mean Well – PN GSM36B15-R7B)
USB cable: connects SphygmoCor XCEL device to PC or laptop. Part number: 1-00858 - USB Cable 2m



Tray (optional) – provides storage of cuffs and tonometer, as well as a mounting platform for the SphygmoCor XCEL device (The storage tray is an optional component supplied by AtCor Medical.)

Part number: 1-00896 - EM4 Storage Tray

Connect Components

Warning: Do not connect any power supply to the SphygmoCor XCEL device other than the power supply provided by AtCor.

- 1. Connect the power supply to the SphygmoCor XCEL device and plug in the power supply to an AC power outlet.
- 2. Connect the SphygmoCor XCEL device to the PC using the USB cable.
- 3. Connect the tonometer to the SphygmoCor XCEL device.
- 4. Connect the femoral cuff to the SphygmoCor XCEL device.
- 5. Power on the SphygmoCor XCEL device.

Note: To disconnect the tonometer, gently push back the connector plug and the connector will automatically release away from the module. DO NOT twist the Tonometer connector.

Start the Measurement

Note: The patient's limb movement should be minimised while capturing data as movement may prevent the capture from being completed or reduce quality of the measurement.

1. Select the Setup screen.

B	Syste	m Database Help				- 🖬	
	Mode PWV			Setu		eport SphygmoCor XCE	
						Height: m Notes:	
Ī	Patient ID	Last Name	First Name	Date Of Birth	Gender	SYS / DIA: / mmHg	
	1	Smith	John	1/1/1954	Male		
	2	Doe	Jane	4/7/1970	Female		
	3	Roe	Jane	10/16/1966	Female	· · · · · · · · · · · · · · · · · · ·	
H	4	Smith	Johnnie	7/8/1998	Male		
						Subtraction method:	
	Patient II First Nam	D: (5) Me: (Janie	Date of Birth	: Dec 30	Hide 1997	Fernoral to Cuff* 200 mm	
	New	Search Edit De	elete Save Cancel			PWV Distance: 355 mm	

2. Select **PWV** mode. (Only required if you have a license for PWA as well).

Syste	em Database Help			
Mode PW	/ •		Setu	o Capture F
PWV PWA	Patients 5			
Patient ID	Last Name	First Name	Date Of Birth	Gender
1	Smith	John	1/1/1954	Male
2	Doe	Jane	4/7/1970	Female
3	Roe	Jane	10/16/1966	Female
4	Smith	Johnnie	7/8/1998	Male
5	Doe	Janie	12/30/1997	Female

Select the patient (by searching Patient ID, Name or Date of Birth, if necessary). If the patient does not exist in the database, create a new patient record by clicking **New** and entering the patient details. Mandatory fields are indicated by *.

					Hide
P	atient ID:			Dat	te of Birth*: Month Day Year
Fin	st Name:	_	_		Gender*:Select
Last	t Name*: <mark>Subject</mark>				
New	Search	Edit	Delete	Save	Cancel

- 3. If the patient's measurement is to be compared to the European General population, the brachial systolic and diastolic pressures should be entered. The brachial pressure obtained from a PWA measurement may be used (if applicable) or from a measurement using a validated sphygmomanometer.
- 4. Position the femoral cuff around the patient's thigh, as high as possible.



5. Measure the PWV Distance

Note: The default setting for the femoral to cuff distance is 200mm. To improve the accuracy of the PWV reading, measure the distance from the top edge of the cuff to the femoral artery, and enter this distance in place of the default. The default setting of 200mm can be changed in the PWV settings menu.

To determine the PWV Distance using the **subtraction method**:

First, measure the *Sternal Notch to Cuff* (top edge of the cuff to the sternal notch) distance in mm using a tape measure, and enter this value into the box marked "Sternal notch to cuff". Typical values are between 400 and 700 mm. Next, measure the distance between the *Carotid and the Sternal Notch* in mm, and enter this value in the box marked **"Carotid to Sternal notch"**.

Enter the measured **"Femoral to cuff distance"**, if applicable. The PWV distance that will be used in the calculation will be displayed below the diagram.



An alternative method of determining the PWV distance is the **direct method**:

Measure the *Carotid to Cuff* (top edge of the cuff to the carotid artery) distance in mm using a tape measure, and enter this value into the box marked "**Carotid to Cuff**". Typical values are between 500 and 800 mm.

Enter the measured "**Femoral to Cuff distance**" if you do not want to use the default value. The PWV distance that will be used in the calculation will be displayed below the diagram.



6. Locate the patient's carotid pulse and position the tip of the tonometer on it. Once the SphygmoCor XCEL system detects a regular carotid pulse, the femoral cuff will be inflated. As soon as the detection period for both the carotid and femoral pulses is complete, the waveforms will turn green and a report will be generated for this measurement. The cuff will then deflate.



Note: The tonometer provides un-calibrated pressure pulse signals. The carotid waveforms displayed are only for qualitative purposes.

If **Cuff/Tonometer Sync** mode is turned off (selected in the PWV settings menu), measure the carotid signal using the tonometer. When the signal quality is valid (the waveform signal colour changes to green), click **Capture** to inflate the femoral cuff and obtain both the carotid and femoral waveforms. The **Capture** button is enabled when there are sufficient signals from the carotid and femoral arterial sites.

Note: For best results, click the Capture button when both waveforms are green.

Once capture is complete, the PWA Report screen is automatically displayed.

Reviewing the PWV Report

System Database Help		- 🛛
Mode PWV 💽 Setup	Capture Report	SphygmoCor XCEL
John Smith Patient Details John Smith Patient ID: 1 Gender: Male Date Of Birth: 1/1/1954 Age: 62 Measurement Details Report Interpretation Height: 178 cm SYS / DIA: 120 / 80 mmHg Notes: PWV Method: Direct PWV Distance: 500 mm	Pulse Wave Veloci Pulse Veloci P	ty Carotid Femoral Ppean General Population (SYS: 120-129 mmHz)
Edit	4 20 35 50 65 80 20 Age (years)	36 50 65 80 Age (years) malised to 80% Direct Distance
Assessments:	Heart Rate Pulse Wave Velocity	Pulse Transit Time QC
6/16/2016 6:06 PM	4 77 bpm 5 8.1 m/s	62 ms 7 🗸
6/16/2016 6:06 PM		
	Delete	Print Patient Report Print
	Simulation data	SphygmoCor

Report data

1 Pulse Wave Velocity – displays femoral and carotid arterial pulses that were recorded during the measurement. The carotid waveform is displayed in blue femoral waveform is displayed in black.

Each waveform has status markers indicating the start of each pulse. A green marker indicates that the pulse was used in the PWV calculation. A red marker indicates that the pulse was not used in the PWV calculation.

2 Healthy Population graph – this shows the patient's overall result in the context of other healthy people of the same age (using data obtained by the University of Cambridge, UK⁵).

Buropean General Population graph – this shows the patient's overall results in the context of a general population that included -patients with diabetes⁶.

⁵ McEniery et al Normal Vascular Aging: Differential Effects on Wave Reflection and Aortic Pulse Wave Velocity: The Anglo-Cardiff Collaborative Trial (ACCT), Am Coll Cardiol 2005;46:1753–60

⁶ Boutouyrie et al, Determinants of pulse wave velocity in healthy people and in the presence of cardiovascular risk factors: 'establishing normal and reference values, European Heart Journal 2010 Oct;31(19):2338-50

4 Heart Rate – average heart rate in beats per minute during the measurement.

5 Pulse Wave Velocity – carotid femoral pulse wave velocity in meters/second (m/s), calculated as the distance divided by the pulse transit time. PWV is averaged over the reading.

6 Pulse Transit Time – time for the pulse to travel from the central arteries to the femoral artery, in milliseconds. The time is measured from the foot (start) of each pulse. The Pulse transit time is averaged over the reading.

QC– Quality Control of the PWV measurement. QC is an indicator of PWV pulse-to-pulse variability over the recording period. High PWV variability indicates a potentially inaccurate PWV measurement as the pulse transit time may not be accurate. An inaccurate measurement is marked with a red cross and the measurement should be repeated. Low PWV variability indicates a high-quality PWV measurement and is shown with a green tick (check mark). The QC indicator should be reviewed after every measurement.

3 Assessments – after data capture is completed, the Report screen is automatically displayed. All measurements are automatically saved and can be viewed at any time by selecting the patient from the Patient screen and pressing the Report button. The most recent report is displayed. Each Measurement is stamped with the date and time of the measurement, with the oldest measurement listed first. If the patient has more than one assessment, clicking on any assessment allows viewing of the corresponding report. A different measurement can also be selected by pressing the arrow key or the tab key on the PC keyboard

Deleting Reports

Each measurement is automatically saved in the database. Reports can be deleted, if desired, as follows:

1. Click **Delete** to remove it from the database.



2. Click Yes.

Note: Reports can be deleted at any time.

Repeat the Measurement

Click the **Repeat** button, which will only be enabled at the last assessment, to perform another measurement on the same patient and create another assessment for the patient.

Return to Patient List

Click **Setup** (at the top of the screen) to return to the list of patients.



Viewing Multiple Reports with the Analysis Screen

Use Analysis to compare multiple sets of data for the selected patient.

- 1. Select PWV mode.
- 2. Click Report.
- 3. Select the first assessment (click on the date and time of the desired assessment).
- 4. Press **Shift** and select another assessment. If it is not adjacent to the first assessment, all of the assessments between these two are also selected.
- 5. The selected assessments are plotted on the screen.

System Database Help					- 🛛
Mode PWV 💌 Setup		Analysis			SphygmoCor XCEL
Patient Details John Smith	1	5	Pulse Wave Velocity		
Patient ID: 1 Gender: Male Date Of Birth: 1/1/1954 Age: 62		D-	•	•	_
Measurement Details Report Interpretation		5 -			
Height: 178 cm SYS / DIA: 120 / 80 mmHg		,	6/16/2016 6:05 PM 6/16/2016 6:06 PM	6/16/2016 6:06 PM	
Notes:		0	Heart Rate		
PWV Distance: 50 cm	Heart Rate (bpm) 2 8	0-	00	•	
		0	6/16/2016 6:05 PM 6/16/2016 6:06 PM	1 6/16/2016 6:06 PM	
Edit					
6/16/2016 6:05 PM		неап кате	Pulse wave velocity	Puise Fransit Lime	QU
6/16/2016 6:06 PM			8.1 m/s		 Image: A second s
	Del	ete			Print

- 6. (Optional) Click **Print** to print a copy of the PWV analysis report.
- 7. Click **Setup** to exit the Analysis screen.

Viewing PWV Reports

You can recall data from any previous session and display it as a report.

- 1. Select the **Report** tab.
- 2. Select the measurement from the Assessments list.

Printing PWV Reports

The **Print** buttons provide a choice of printed reports.

Print Patient Report Print

Print – a single-page PWV report.

Print Patient Report – a multi-page report that provides an explanation of the different PWA

parameters. This report may not be available in some regions.

Printed PWV Report Example



AtCor Medical SphygmoCor XCEL 1.3.2.10 9/26/2016 12:37 PM

Automatic versus Manual Waveform Capture

Carotid Waveform capture can be performed either automatically or manually.

- If Auto Capture is enabled (default setting), the software will automatically capture the waveform when all of the quality parameters are met. Alternatively, the keyboard space bar can be pressed at any time to override the Auto Capture feature and capture the carotid waveform.
- If Auto Capture is disabled, data capture can be completed by manually pressing the spacebar on the computer keyboard when the minimum number of reliable pulses has been recorded and the capture button has become illuminated.

Using the Guidance Bars

SphygmoCor XCEL System provides feedback (guidance bars above and below the waveform) to aid in determining if the waveforms are consistent enough to provide an accurate PWV measurement.

As shown in the following diagram, there are multiple visual indications that provide feedback.

If Auto Capture is used (default setting), the data is captured automatically and calculation is performed to produce a report after the waveform colour changes to green. This indicates quality waveforms for the duration of the capture time. For PWV, both carotid tonometer and thigh cuff signals need to be green for the data to be automatically captured and a valid report produced. Data capture can be initiated at any time by clicking the **Capture** button, regardless of the waveform colour.



Waveform – the waveform changes colour according to the quality of the overall recording during capture time. Initially, the waveform is white and changes to green if the quality of the overall recording during the capture time is acceptable and the recording has at least the minimum number of quality pulses. This number varies according to the data capture duration selected in setup.



2 Top guide bar – displayed at the top of the waveform, indicating reliability of the pulse peaks. It updates beat-by-beat. It is green if acceptable for data capture and red if not recommended for data capture. For the tonometer carotid signal only, it can also be yellow, indicating marginal data.

If the top guide bar of the tonometer signal is not green, apply more pressure to the tonometer. If that is not successful, try moving the tonometer slightly.

3Bottom guide bar – displayed at the bottom of the waveform, indicating reliability of the pulse troughs. It is green if acceptable for data capture and red if not recommended for data capture. For the tonometer only, it can also be yellow, indicating marginal data. Apply a consistent pressure on the tonometer to ensure that the signal remains green.

4 Bar graphs – indicate signal amplitude, measured beat-by-beat. The amplitude must be above the threshold mark **5** to ensure a reliable signal.

The waveform colour is synchronised with the bar graph colour:

- when the waveform is green in colour, the bar graph is also green
- when waveform is white in colour, the bar graph is red in colour

Note: Guidance Bars and Auto Capture are part of the Capture Guide feature in the software. The default setting includes enabling of both Auto Capture and Guidance Bars. These features can be enabled or disabled from the System menu (refer to

PWV Settings)

Techniques for Obtaining Quality PWV Measurements

Fitting the femoral cuff

Correct fitting of the cuff helps to ensure a reliable reading. Use the following pictures as a guide.

Note: To ensure reliable readings, the thigh cuff should be placed over bare skin, or over thin clothing or material.

1. Place the cuff in position around the patient's upper thigh, with the tube on top of the leg and facing upwards (towards the patient's head).



2. Wrap the cuff firmly but not too tight.



3. Ensure the cuff fits neatly, as shown in this picture.



Control Conditions

When performing a SphygmoCor XCEL PWV measurement, it is recommended that the patient should:

- Abstain from alcohol for at least 6 hours prior to the measurement.
- Abstain from tobacco and caffeine for at least 4 hours prior to the measurement.
- Fast for at least 6 hours prior to the measurement, however, if this cannot be achieved, a light meal before the measurement is allowed.
- Rest (seated or supine position) for at least 5 minutes prior to the measurement.
- Be as relaxed as possible and not talk during the measurement.
- When possible, subsequent measurements should be obtained at the same time of the day so as to account for the diurnal blood pressure effect.

Tonometer Technique

Placement of the tonometer is important to ensure quality waveforms:

- Remove the tonometer from either the tonometer storage tray or the temporary holder on the top of the SphygmoCor XCEL device. Hold the tonometer base gently but firmly between the tip of the thumb and the fingers.
- Locate the patient's carotid artery by palpation, and identify the strongest pulse point.
- Press the tonometer gently but firmly against the patient's neck at the strongest pulse point.
- Maintain steady pressure with the tonometer.
- If the trace is flat at the top of the screen, reduce the pressure. If the trace is flat across the bottom of the screen, increase the pressure.



- While holding the tonometer steadily on the patient's neck, the waveform signal should automatically resize with optimal placement. Make minor adjustments until the waveforms are uniform in shape and height.
- When the Guidance Bars are enabled, red, green or yellow bars will appear on the screen at the top, bottom and sides of the waveforms travelling across the screen.
- The Guidance Bars indicate the amount of variation of the waveforms. Red indicates too much variation; yellow indicates moderate variation and green indicates an acceptable level of variation. Messages will appear at the bottom of the screen indicating the area of variation to guide in adjusting the position of the tonometer. Adjust the tonometer slightly medially or laterally over the artery to obtain a clear, strong signal. The following image shows an example of good signals.



System Installation and Setup

Unpacking

Carefully un-pack the SphygmoCor XCEL System.

Remove the SphygmoCor XCEL device from the packaging then remove the protective plastic film.

A smaller carton contains all the accessories included with the system.

Note: AtCor Medical strongly recommends that the SphygmoCor XCEL System is installed on a dedicated computer or notebook computer because some 3rd party software applications may interfere with the operation of SphygmoCor XCEL software.

Software Download

The SphygmoCor XCEL software installation media can be downloaded from the AtCor Medical website.

- 13. Open atcormedical.com in your web browser.
- 14. Navigate to the **Customer Support** section.
- 15. Create an account.
- 16. You will now have access to the XCEL Software, User Manual and Inservice videos.

- 17. To download the required files, right click on each file and select 'Save link as'. Save the files to your local drive.
- 18. To open the XCEL Software zip file, right click on the zip file and extract all files to your PC.

Software Installation

The SphygmoCor XCEL software installation media contains the software needed to install the SphygmoCor XCEL software on a dedicated computer. Complete the software installation prior to connecting the module to the computer.

Note: Administrator privileges are required for installation.

Note: It is recommended to connect the computer to the internet and download and install all Windows Updates before installing the software.

Note: For non-English Windows 8 and Windows 10, before installation users need to add English in the **Region and Language** section under the **Time and Language** section in the **PC Settings**, then download and install the English language package. Please refer to Windows manual for further details.

Note: XCEL 1.3 can be installed on computers that have a previous version of XCEL installed. If SphygmoCor XCEL 1.3 is already installed, go to *Uninstallation* to uninstall the software and install again or go to *Repair* to reinstall features already installed. It is highly recommended to back up your existing database before uninstallation or repair to prevent data loss.

Note: When Installing XCEL 1.3, any available compatible user settings from a previous XCEL installation will be copied across to the XCEL 1.3 settings.

Select the installation media in your computer and follow the Standard Installation if auto play is enabled on your computer. If auto play is not enabled, navigate to the installation media and double click on the setup.exe file \mathscr{D} .

Note: For custom installation, refer to *Custom Installation* section.

Standard Installation

1. Run setup.exe to commence installation.

2. If the following screen appears, click Install, otherwise go to 2.

S	SphygmoCor XCEL Software Suite - InstallShield Wizard				
	🎸 🖁	he SphygmoCor XCEL Software Suite requires the following items to be installed on your omputer. Click Install to begin installing these prerequisites.			
	Status	Requirement			
	Pending Pending	Microsoft .NET Framework 3.5 SP1 (Windows Feature) Microsoft .NET Framework 4.5.1 Full			
		Install Cancel			

3. Close all other programs and click **Install**.

XCEL Installer	×
	Welcome to XCEL Installer. This program will install the SphygmoCor XCEL Software Suite on your computer. Please exit all other programs then click Install. WARNING: This program is protected by copyright law and international treaties. PhygmoCor Cor Medical
XCEL: 1.3.1.4 Installer: 130.1.0.1 SN: XCEL_1.3 (66666).1234567890	Install Exit

4. Read the license agreement and if you agree to the terms check the box **I accept the terms of the license agreement** then click **Next**.





5. The default set up type is Complete and the default destination folder is C:\AtCor\XCEL 1.3. Keep the default settings and click **Next**. (For Custom installation refer to *Custom Installation* Section to continue installation)

XCEL Installer	3
Select a setup type and install destination folder that best suits your needs.	
Please select a setup type. Complete All features will be automatically configured and installed locally. Third-party software license agreements will be accepted on your behalf.	
Custom	
Destination Folder C:\AtCor\XCEL 1.3 Browse TostallShield	
< <u>B</u> ack Next > Cancel]

6. Select the language you would like SphygmoCor XCEL to be displayed in, and then click Next.



Select a language Select a language you would like SphygmoCor XCEL to be in. Select the language you would like SphygmoCor XCEL to be in.	CEL Installer			×
Select the language you would like SphygmoCor XCEL to be in. English (UK) English (US) French (Français) Japanese (日本語)	Select a language		Sphygmo XC SIMPLY THE GOLD STAT	
✓ English (UK) □ English (US) □ French (Français) □ Japanese (日本語)	Select the language you would like SphygmoC	or XCEL to be in		
	<mark>▽ English (UK)</mark> □ English (US) □ French (Français) □ Japanese (日本語)			
	estallChield			
allShield	istalionielu			

Note: The choice of languages might be different from the given list shown in the above image.

7. Click Anyone who uses this computer (all users) to create XCEL shortcuts for anyone who can log in to the computer, otherwise select Only for me, then check the box Check to Create a Desktop Shortcut for SphygmoCor XCEL and click Install.



8. Click Finish to exit and restart your computer if prompted

XCEL Installer	
	Installation Complete XCEL Installer has successfully installed the SphygmoCor XCEL Software Suite. Thank you for your purchase. Click Finish to exit.

Note: If the Installation Complete confirmation message is not displayed, contact AtCor Medical.

Note: A text file report is created as a record of XCEL 1.3 installation. To see the report go to the **XCEL** 1.3 installation folder \rightarrow system \rightarrow logs \rightarrow XCEL Installation Report <timestamp>.txt where <timestamp> is the date and time the file was created.

Custom Installation

For Custom Installation, the operator must go through Step1 to 4 of the *Standard Installation* and then continue with the following steps:



1. Select Custom, and select a destination folder (default C:\AtCor\XCEL 1.3) and click Next.

XCEL Installer		×
Setup Type & D Select a setup t suits your need	Destination Folder Sphyce type and install destination folder that best Is. SIMPLY THE C	SOLD STANDARD
Please select a	setup type.	
Complete		
	All features will be automatically configured and installed lo software license agreements will be accepted on your beha	cally. Third-party alf.
Custom	Allows selection and configuration of features.	
Destination F	older	
C: \AtCor\XCE	£ 1.3	Browse
InstallShield		
	< <u>B</u> ack <u>N</u> ext >	Cancel

2. Select the features you want to install and click Next.





3. Select the language you would like SphygmoCor XCEL to be displayed in, and then click **Next**.

XCEL Installer				×
Select a language			Sphygmo XC SIMPLY THE GOLD ST/	
Select the language you w	ould like SphygmoCo	r XCEL to be in.		
<mark>☑ English (UK)</mark> □ English (US) □ French (Français) □ Japanese (日本語)				
nstallShield		< <u>B</u> ack	<u>N</u> ext >	Cancel

Note: The choice of languages might be different from the given list shown in the above image.

4. Click **Anyone who uses this computer (all users)** to create XCEL accessible to anyone who can log in to the computer, or select **Only for me,** then check the box **Check to Create a Desktop Shortcut for SphygmoCor XCEL** and click **Install**.

XCEL Installer			
Ready to Install XCEL Installer is ready to begin installation. SphygmoCor XCEL SIMPLY THE GOLD STANDARD			
Click Install to begin the installation.			
If you would like to review or change any of your installation settings, click Back. Click Cancel to exit.			
Install SphygmoCor XCEL for:			
 Anyone who uses this computer (all users) 			
Only for <u>m</u> e (parisak)			
Create a Desktop <u>S</u> hortcut for SphygmoCor XCEL			
InstallShield			
< <u>B</u> ack Install Cancel			

5. Click **Finish** to exit and restart your computer if prompted.

XCEL Installer	
	Installation Complete XCEL Installer has successfully installed the SphygmoCor XCEL Software Suite. Thank you for your purchase. Click Finish to exit.
	Finish

Note: If the Installation Complete confirmation message is not displayed, contact AtCor Medical.

Note: Custom Installation can be used to setup SphygmoCor XCEL in a network configuration. For more information on network configuration using custom installation refer to the *Appendix* section.

Uninstallation and Repair

If you want to uninstall the SphygmoCor XCEL 1.3 software go to *Uninstallation*. Otherwise, go to *Repair* to reinstall features previously installed.

Note: It is recommended to back up your database before uninstallation.

Uninstallation

To uninstall the XCEL 1.3 software, you have two options: Direct Uninstallation or Maintenance Mode.

a) Direct Uninstallation

Go to **Control Panel→ Programs→ Programs and Features** and uninstall the SphygmoCor XCEL 1.3 program.

b) Maintenance Mode

Select the SphygmoCor XCEL installation media and run setup.exe. The installer will automatically go to Maintenance mode if the SphygmoCor XCEL 1.3 software is already installed.

1. Select Remove in and click Next to continue.



2. Click Yes to confirm uninstallation.

XCEL Installer - Maintenance
Are you sure you want to uninstall the SphygmoCor XCEL software? The Atcor database will not be uninstalled.
Yes <u>N</u> o

3. Wait until the Uninstall Complete confirmation will appear. Click Finish to exit.



Repair

Select the SphygmoCor XCEL installation media and run Setup.exe. The installer will automatically go to Maintenance mode if the SphygmoCor XCEL 1.3 software is already installed.

1. In the Maintenance Welcome screen, select **Repair** and click **Next** to reinstall the features you have already installed.



2. Wait until Maintenance Complete screen appears. Click Finish to exit.



Note: When running installer in the Repair mode, current XCEL settings will be reset to default settings of the previously installed language. The password set in the Clinical Trials Option will also be cleared. To set a new password refer to *General Settings* section.

Deleting or Importing SphygmoCor CvMS Database

If SphygmoCor CvMS version 8 or version 9 is installed on your computer, the SphygmoCor XCEL1.3 software will copy the CvMS database file from SphygmoCor CvMS version 8 or version 9. When the operator runs SphygmoCor XCEL1.3 for the first time, the following message will appear for the operator to confirm the migration of patient's records from the CvMS database to the current XCEL1.3 database.



Click **Yes** if you would like to accept the migration. By clicking **No**, the message will appear the next time the user starts XCEL software. To avoid migration of the CvMS database in future, the CvMS database files must be deleted from the XCEL directory below.

Deleting SphygmoCor CvMS Database

To delete the CvMS database files follow these instructions:

- 1. Go XCEL installation directory e.g. Local Disk (C:) \rightarrow AtCor \rightarrow XCEL 1.3 \rightarrow system \rightarrow data
- 2. Delete the CvMS database file (.xyz or .mdb file extension)

Importing SphygmoCor CvMS Database

To import patient details from a CvMS database file manually follow the procedure below. The CvMS database files must be in .mdb or .xyz format.

- SphygmoCor CvMS V8
- 1. Go to the SphygmoCor CvMS installation directory- Local Disk (C:) → Program Files (x86)/ Program Files → AtCor → SphygmoCor CvMS → data
- 2. Copy the CvMS database (.xyz or .mdb file extension) into the **XCEL installation directory Local Disk (C:)** → **AtCor** → **XCEL 1.3** → **system**→ **data**
- SphygmoCor CvMS V9
- 1. Go to the SphygmoCor CvMS V9 installation directory Local Disk (C:) → AtCor → SphygmoCor CvMS V9 → data
- 2. Copy the CvMS database (.xyz file extension) to XCEL installation directory Local Disk (C:) → AtCor → XCEL 1.3 → system→ data

Starting SphygmoCor XCEL software

- 1. Ensure the SphygmoCor XCEL device is set up as described in the *Connecting Components* section.
- 2. Connect the SphygmoCor XCEL device to the computer and power the device on using the switch on the rear of the device. The green power and orange standby indicators will illuminate.



- 3. SphygmoCor XCEL software can be started in either of two ways:
 - Select Start → All Programs → SphygmoCor XCEL 1.3 → 💅 XCEL 1.3.
 - Double-click the XCEL 1.3 software icon 💉 on the desktop, if the shortcut is created in the installation process.
- 4. Enter System Key (as provided in the System Information Leaflet).

🎸 System Key		×
Please enter a va	alid System Key	
		· · ·
	Oł	Cancel

Note: If SQL Server 2014 Service is in the system and not running, XCEL will start SQL Server 2014 Service. Therefore Windows system will pop up an UAC notification message depending on the user account type:

Administrator account

Click Yes, XCEL will start SQL Server 2014 Service and it will continue to run as normal.

Survey User Account Control	\times
Do you want to allow this app to make changes to you PC?	r
Program name: Net Command Verified publisher: Microsoft Windows	
Show details Yes No	
Change when these notifications app	<u>ear</u>

Users account

Enter administrator **Username** and/or **Password** if one of the following messages is displayed. Click **Yes**, XCEL will start SQL Server 2014 Service and run as normal.



۲	User Account Control	×
Do yo chan	ou want to allow the following program to ges to this computer?	make
	Program name: Net Command Verified publisher: Microsoft Windows File origin: Hard drive on this computer	
To continue	, type an administrator password, then click Yes.	
P	Username Password Domain:	
Show det	ails Yes	No
۲	User Account Control	×
Do ye chan	ou want to allow the following program t ges to this computer?	o make
	Program name: Net Command Verified publisher: Microsoft Windows File origin: Hard drive on this computer	
To continue	, type an administrator password, then click Yes.	
P	Admin-Username	

- 5. In the Setup screen on the top right, click on System menu and select Find Module
- 6. The message *Electronic Module Found* appears at the bottom of the **Setup** screen indicating that the PC has identified the connection to the SphygmoCor XCEL device.

Note: If the message does not appear, check the USB connection and confirm that the SphygmoCor XCEL device is powered ON. Repeat Step 4. If the problem persists, please contact AtCor Medical for technical support.

7. The SphygmoCor XCEL system is ready.

System Settings

The SphygmoCor XCEL System Settings enable the operator to configure many features and options in the software. The System Settings contains the following:

- 1. General Settings
- 2. PWV Settings
- 3. PWA Settings
- 4. Brachial BP Settings

General Settings

General PWV PWA Brachial BF	×
Startup Mode PWV PWA Setup Screen Patient Privacy Height Unit Metric Imperial Comms Port Simulation	Reports Report Title (AtCor Medical SphygmoCor XC) Report Logo Browse Extra Features Clinical Trials Option
	Save Cancel Default

Startup Mode – PWV / PWA – this setting determines the default mode when the SphygmoCor XCEL software is started (if both modes are enabled in the software). Click on PWA or PWV to change the setting. The selected start up mode is marked with a green dot.

Setup Screen – Patient Privacy – this features offers the ability to temporarily hide the list of patients to maintain patient privacy. A check mark appears in the box when patient privacy has been selected.

Height Units – SphygmoCor XCEL software can display height in either metric (cm) or imperial (feet/inches) measurements.

Comms Port – This setting identifies which communications (comms) port on the PC is connected to the module as well as enables simulation of SphygmoCor data for demonstration purposes. If Simulation is selected, the software will display demonstration data and *not* actual patient data. When obtaining a measurement on a patient, the communications port (eg, com3, com4, etc) should be displayed. The Simulation setting should only be chosen when demonstrating the software functionality *without* a patient.

Reports – Report Title, Report Logo – This section enables customization of the title on printed reports, eg, hospital or clinic name, as well as incorporation of customised logo on the printed report. To modify the report name, move the cursor to the space under "Report Title" and enter the title you would like printed on reports. To use your organization's logo on reports, the logo must be stored on the computer (as a .jpg or .bmp file). Click "Browse" and select your logo from the appropriate file. It is recommended to import a 512 × 512 pixel logo image for maximum resolution.

Extra Features – Clinical Trials Option – When Clinical Trials Option is selected, a **Clinical Trials Option** tab will appear in the settings allowing the appropriate personnel to define descriptions for individual visits of a clinical trial and to protect the settings by a password.

Note: Simulation Mode will not be enabled if the Clinical Trials Option has been selected.


General PWV PWA Brachial	BP Clinical Trials Option
Startup Mode PWV PWA Setup Screen Patient Privacy Height Unit Metric Imperial Comms Port <u>COM1</u>	Reports Report Title AtCor Medical SphygmoCor XCI Report Logo Browse Extra Features Clinical Trials Option
	Save Cancel Default

When **Clinical Trials Option** is selected, two fields of **Operator** and **Visit** will appear; for the operator's name and patient's visit description respectively in the PWA and PWV Setup and Report screen.



PWA Setup



PWV Setup

In the **Report** Screen, the operator's name and patient's visit description will appear in the *Measurement Details* section of PWA Report and PWV Report as following:



	PWA Report
Patient Details Patient ID: Date Of Birth:	John Smith 1 Gender: Male 12/1/1900 Age: 115
Measurement Details	Report Interpretation
Brachial SYS/DIA: Height: Operator:	160 / 120 mmHg QC: ✓ ➡ 188 cm
Notes:	

PWV Report					
Patient Details John Smith					
Patient ID: 1 Gender Date Of Birth: 1/1/1954 Age	Male 62				
Measurement Details Report Interpretation					
Height: 178 cm Operator					
SYS / DIA: 120 / 80 mmHg Visit					
PWV Method: Direct PWV Distance: 50 cm					

Click on the **Clinical Trials Option** heading tab in the Settings window to define visit descriptions and set the optional password to protect the settings:

a) Visit Description – Click on the first row to add a visit description. To edit a visit description, double click on the existing visit description. To delete a visit description, click on the visit description, and press **Delete** button on the keyboard.



	VV PVVA Brachi		
	Vis	it Description	
	Click her	e to add a new row	
Visit One: fi	rst time		
Visit Two: s	econd time		
Visit Three:	third time		
Visit Four: f	ourth time		
Visit Five: fi	fth time		
Visit Six: six	th time		
Visit Seven:	seventh time		
		Confirmation:	
	Protect		

Note: In **Clinical Trials Option**, when a new patient is created, the patient "Last Name" field is automatically populated with the text "Subject".

- b) **Password** The appropriate personnel can select a password to protect **Settings** from other users. The correct password is required to change any of the settings.
 - To Protect the Settings with a password:
 - 1. Go to System → Settings → General → Extra Features
 - 2. Check Clinical Trials Option box in Extra Features (a new window will be opened)
 - 3. Go to Clinical Trials Option.
 - 4. Enter your password in the **Password** field.
 - 5. Re-enter the selected password in the **Confirmation** field.
 - 6. Click **Protect** to lock the settings.
 - 7. Click **Save** to save changes.

	Visit	Description	
	Click here	to add a new row	
Visit One: first f	time		
Visit Two: seco	nd time		
Visit Three: thir	d time		
Visit Four: fourt	th time		
Visit Five: fifth	time		
Visit Six: sixth t	ime		
Visit Seven: se	venth time		-
	Password:	Confirmation:	
*	****	*****	
	Protect		

- To Unprotect the Settings using the selected password:
 - 1. Go to System → Settings → Clinical Trials Option
 - 2. Enter the password in the **Password** window.
 - 3. Click **Unprotect** to unlock the settings.
 - 4. Click **Save** to save changes.

	Visit Description	on		<u></u>
	k here to add a r			
Visit One: first time				
Visit Two: second time				
Visit Three: third time				
Visit Four: fourth time				
Visit Five: fifth time				
Visit Six: sixth time				
Visit Seven: seventh time				
Password: ********* Unpre	tect			

Note: If the wrong password is entered the **Invalid Password** warning will appear. If the selected password is forgotten, go to the Repair mode in *Repair* section to reset the password.

PWV Settings

These settings are only available if PWV software was purchased with the SphygmoCor XCEL System.

4	Settings	x
	General PWV PWA Brachial Bl	P
	Capture Time 5 seconds 10 seconds 20 seconds Capture Guide Guidance Bars Auto Capture	PWV Distance Unit mm cm Femoral to Cuff Distance 200 mm Report Screen
	Subtraction (sternal notch to cuff)	 sternal notch to carotid)
		Save Cancel Default

Capture Time – this setting refers to the length of time for which a stable reading must be available for a successful PWV measurement. The default setting is 10 seconds. For patients with a stable heart rate, the setting can be changed to five seconds. For patients with unstable heart rates, the 20 second capture may be required to obtain a valid measurement.

Capture Guide features

- Guidance Bars this setting enables visual guides that give feedback of the signal quality from both tonometer and cuff while a measurement is in progress. Refer to *Using the Guidance Bars* for an explanation of how Guidance Bars can improve data capture. The default setting is Guidance Bars enabled.
- Auto Capture this setting results in automatic capture of the recording as soon as reliable tonometer and cuff signals are detected simultaneously. If not selected, capture must be manually triggered by pressing the space bar (Manual capture can be achieved by pressing the space bar, even if Auto Capture is enabled). Default setting is that Auto capture is enabled.
- Cuff/Tonometer Sync mode this setting causes the thigh cuff to inflate when a reliable carotid signal from the tonometer is detected. If disabled, the cuff inflation is triggered as soon as the capture starts. The default setting is disabled.



Simulation Data – this setting offers the ability to demonstrate PWV simulation mode (i.e., without a patient). The PWV data file contains both carotid tonometer and thigh cuff signals. Switching between real measured data and simulated data is controlled by the COM Port setting under *General Settings*. Actual patient data will not be obtained when the Simulation is enabled.

PWV Distance Method – this setting determines how the SphygmoCor XCEL System calculates the distance between the patient's carotid measurement site and the cuff. There are two commonly used methods supported by SphygmoCor XCEL.

- Direct method the distance is measured directly from the carotid artery to the top edge of the femoral cuff.
- Subtraction method distance is measured from the sternal notch to the top edge of the femoral (thigh) cuff (distal distance), and from the carotid artery to the sterna notch (proximal distance). The proximal distance is subtracted from distal distance to provide the aortic distance, which is used in the calculation of PWV. The subtraction method is the default setting.



PWV Distance Unit – this setting offers use of either centimetres or millimetres as the PWV distance units. Millimetres (mm) are the default option.

Femoral to Cuff Distance – a default distance between the top of the cuff and the femoral artery of 200mm is used during PWV measurements (PWV Setup Screen). The value can be changed in the PWV Set up screen.

Report Screen – the PWV report will include reference PWV values from the European General Population to be compared with the measured PWV if this option is selected. This reference population includes healthy subjects and subjects with cardiovascular risk factors. The default setting includes the European General Population data on the report.

PWA Settings

These settings are only available if PWA software was purchased with the SphygmoCor XCEL system.



🎸 Settings	x
General PWV PWA Brachial BP	
Capture Time 5 seconds 10 seconds 20 seconds	Capture Input Pressure Cuff Tonometer
 Capture Guide ✓ Guidance Bars ✓ Auto Capture 	● Quick Start ● Standard
Simulation Data	
	Save Cancel Default

Capture Time – this setting refers to the length of time for which waveforms will be collected for a successful PWA measurement. A higher capture time may be needed for patients with slow respiratory cycles and/or a highly variable heart rate. The default setting is 5 seconds but for patients with slow respiratory cycles and/or a highly variable heart rate, a higher capture time may be needed for a valid measurement.

Capture Guide

- Guidance Bars if selected, colour-coded Guide Bars are displayed, indicating the quality of recorded pulses. Refer to *Using the Guidance Bars* for an explanation of how Guidance Bars can improve data capture. The default setting is Guidance Bars enabled.
- Auto Capture if selected, automatic capture of the recording will occur once reliable quality cuff signals are detected ie, colour is green. If not selected, capture must be triggered manually by pressing the keyboard space bar. Refer to the *Automatic versus Manual Waveform Capture* section under PWA. The default setting is Auto Capture enabled.

Simulation Data – this setting offers the ability to demonstrate the PWA software using simulated data (i.e., actual patient data will not be captured when this Simulation is selected). Simulation must also be selected under *General Settings* in order to use the "Demo" setting.

Capture Input – this setting determines whether the signal is captured from the Pressure Cuff (i.e. brachial upper arm cuff) or radial artery tonometer. For SphygmoCor XCEL software version 1, Pressure Cuff is the only available option.



Brachial BP Settings

🎸 Settings 🛛 🐥
General PWV PWA Brachial BP
Auto PWA ✓ If SYS >= 0 ✓ If DIA >= 0
✓ If PP >= 0
Assessments
Number of Assessments
Initial Rest 0 seconds
Subsequent Rest 5 seconds
Save Cancel Default

Auto PWA – this setting enables threshold values for brachial systolic, diastolic or pulse pressure (SP, DP and PP respectively). If the measured blood pressure is above the threshold, PWA waveform capture commences automatically. If the measured Brachial Blood pressure is below the threshold or all check boxes are not selected, the PWA waveform assessment will only be performed if **Capture** is clicked.

Note: It is recommended that the user click **Capture** as soon as the brachial BP measurement has ended and the brachial BP values are displayed. Also, it is recommended that the patient refrain from movement and remain in a stable position until the PWA measurement ends.

When enabled, this option can be used to screen patients with potential cardiovascular risk indicated by high brachial blood pressure. The default setting is 0 for each, thus the PWA waveform capture always starts automatically.

Number of assessments – this setting enables the captured waveform to be calibrated with multiple readings of brachial systolic and diastolic cuff pressure. The number of brachial assessments can be set from 1 to 5, with 1 being the default value. Studies have shown that the average of multiple measurements of brachial blood pressure may provide a more accurate measure than one individual brachial blood pressure measurement.⁷ The operator can also choose to ignore the first brachial BP measurement by ticking the **Ignore the First Assessment** checkbox.

⁷ Vidt, DG et al. Taking Blood Pressure: Too Important to Leave to Humans? Cleve Clin J Med. 2010 Oct; 77(10):683-8

Initial Rest – Allow the operator to choose the time to start the first brachial BP assessment after clicking the **Start** button. The default initial rest is 0 seconds.

Subsequent Rest – Allow the operator to choose the time between subsequent brachial BP assessments when the operator chooses more than one brachial BP assessment. The default (and minimum) subsequent rest is 5 seconds.

SphygmoCor XCEL Device Status Indications

Indicators on the SphygmoCor XCEL device indicate the device status, as shown in the following diagram and table.







Number	Name & Symbol	Colour and state	Status
	Power	Off	Power is off
1	С	Green	Power is on
	Status	Yellow	Software is initializing
		Flashing Yellow	Software initialization failure
2	ίΟ ^ν	Flashing Red or off	Device requires service (Return to manufacturer)
		Flashing Green	Communicating with PC
		Green	Operating normally (waiting)
	Cuff	Blue	Active
3	Ò	Off	Any other condition
	Tonometer	Blue	Active
4	ſ	Off	Any other condition
6	Stop	Red	STOP button had been pressed and the cuff is deflating.
		Off	Operating normally (cuff ready to inflate)
		Flashing Red	Safety cuff deflation by device

Creating and Managing Patient Records

Before creating a new patient entry, confirm that the patient's information has not already been entered by searching the database as separate patient entries cannot be merged.

To locate a patient record

- 1. Click **Search**. The patient fields become white.
- 2. Enter the known patient details into any of the search fields (Patient ID, First Name, Last Name, Date of Birth, and Gender).



Example: enter **s** in the first name field.

Note: the software will begin searching as soon as data is entered in any searchable field. The set of results becomes smaller as more data is entered.

Patients 5					
Patient ID	Last Name	First Name	Date Of Birth	Gender	
2	Doe	Jane	4/7/1970	Female	
5	Doe	Janie	12/30/1997	Female	
3	Roe	Jane	10/16/1966	Female	
1	Smith	John	1/1/1954	Male	
4	Smith	Johnnie	7/8/1998	Male	

- 3. Click any of the column headings to sort the list of results according to data in that column, such as Last Name. Click the heading again to sort in reverse order.
- 4. Click anywhere in the patient's record to select that patient.
- 5. Click **Cancel** to end the search and return to the full list of patients.

To create a new patient record:

1. Click New.

The patient entry fields will become white, indicating the software is ready for new patient entry.

					l	Hide
Р	Patient ID:			Date o	f Birth*: Month Day	v ▼ Year ▼
Fir	rst Name:	_		G	ender*:Select	
Las	st Name*: <mark>Subjec</mark>	t				
New	Search	Edit	Delete	Save	Cancel	

2. Enter all mandatory data, indicated by the * symbol. Other data may also be entered but is optional.

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3. Click Save.

Editing an Existing Patient

To edit a record:

1. Select the record (search, if necessary).

	Hide
Patient ID: (5	Date of Birth*: Dec 30 1997
First Name: (Janie	Gender*: (Female
Last Name*: Doe	
New Search Edit Delete Save	Cancel

Initially, the record is read-only, which is shown by white text.

2. Click Edit.

		Hide
Patient ID:	5	Date of Birth*: Dec 30 1997
First Name:	Janie	Gender*: Female
Last Name*:	Doe	
New	earch Edit Delete Save	Cancel

The text changes to black and you can make changes to all fields.

Note: Editing a patient's ID, First Name or Last Name is not case sensitive. If you want to change these fields by uppercase or lowercase, make sure to change the spelling first and save it. Then, modify the field to desired case and original spelling.

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Deleting a Patient

Caution: Deleted data is permanently removed from the database and cannot be retrieved.

To delete a record:

1. Select the record (search, if necessary).

	Hide
Patient ID: 5	Date of Birth*: Dec 30 1997
First Name: (Janie	Gender*: (Female
Last Name*: Doe	
New Search Edit Delete Save	Cancel

To begin with, the record is read-only, which is shown by white text.

- 2. Click Delete.
- 3. Select **Yes** from the confirmation pop-up window to confirm deletion.

Troubleshooting

Caution: If any error messages indicate "database corruption" or "database access error" contact AtCor Medical Technical Support immediately.

The SphygmoCor XCEL Software conducts a series of internal tests with prompts to guide the user should an error occur. In most instances, errors can be resolved following these prompts. However, if the error is not resolved and the solution is not apparent, please contact AtCor Medical Technical Support for assistance.

SphygmoCor XCEL Device

Condition 1: The SphygmoCor XCEL device cannot be detected.

This error may appear at two places in the software:

- When the SphygmoCor XCEL software is started.
- Upon entry into the Capture Screen.

Check the following items:

- The SphygmoCor XCEL device is connected to your computer.
- The SphygmoCor XCEL device is plugged into an AC outlet and the power switch is toggled to the "on" position.
- The Power light is illuminated (if it is not illuminated, refer to Condition 2 "The POWER light is off" below)
- The correct communications port is selected in the configuration settings.

From the Setup Screen, select System – Find Module.

• If using a USB Adaptor, ensure that appropriate USB adaptor drivers have been installed on the computer.

If none of the above actions resolves the issue, please contact AtCor Medical Technical Support.

Condition 2: The POWER light is off.

Check the following:

- The SphygmoCor XCEL device power cable is connected to an AC outlet and the toggle switch is in the "ON" position.
- If the above action does not resolve the issue, please contact AtCor Medical Technical Support.

Software Screens

The user will be notified of software issues via pop-up windows with instructions to resolve the issue. If the screen freezes or you require further explanation of any pop-up window, please contact AtCor Medical Technical Support for assistance.



Capture Screen

The Tonometer fails to respond.

Ensure that the Tonometer is connected to the appropriate port at the rear of the SphygmoCor XCEL device.

Note: Use caution when connecting the Tonometer as pins inside the connector can be easily bent or broken.

Error Codes

The following is a list of possible errors that may be encountered.

Note: If any error occurred where the code is not listed in the following table, exit XCEL software, turn OFF/ON the device and then start XCEL software. If problem persists please contact AtCor Medical Technical support.

Error Code	Error Message	Action
1	Unable to continue Brachial Assessment due to weak oscillometric signal (Code = 1).	 Ensure use of correct cuff size, correct positioning and fit Ensure that no obstructions (eg, clothing) are located between the cuff and the patient's arm After completing these steps, attempt another measurement
2	Unable to continue Brachial Assessment due to erratic oscillometric signal (Code = 2).	 Ensure that the patient's arm is stationary during the measurement Ensure use of correct cuff size, correct positioning and fit After completing these steps, attempt another measurement
3	Unable to continue Brachial Assessment due to exceeded retry count (Code = 3).	 Ensure that the patient's arm remains stationary during the measurement Ensure use of correct cuff size, correct positioning and fit Remove any obstructions between the cuff and the arm After completing these steps, attempt another measurement
4	Unable to continue Brachial Assessment due to exceeded measurement time limit (Code = 4).	 Before retrying the measurement: Ensure that the patient's arm remains stationary during measurement Ensure use of correct cuff size, correct positioning and fit Remove any obstructions between the cuff and patient's arm After completing the above steps, attempt another measurement

Error Code	Error Message	Action
85	Unable to continue Brachial Assessment due to pneumatic blockage (Code = 85).	 Ensure that the hose is not bent or pinched Ensure that the hose and cuff is not under the patient Ensure use of correct cuff size, correct positioning and fit After completing the above steps, attempt another measurement
87	Unable to continue Brachial Assessment due to inflate timeout (Code = 87).	 Ensure that the hose connections to the device and cuff are intact and tight Ensure use of correct cuff size, correct positioning and fit Ensure that the cuff is not leaking air After completing the above steps, attempt another measurement
88	Unable to continue Brachial Assessment due to safety timeout (Code = 88).	 Turn the device OFF and then turn it ON Ensure use of correct cuff size, correct positioning and fit Ensure that the patient's arm is stationary during the measurement After completing the above steps, attempt another measurement
89	Unable to continue Brachial Assessment due to cuff overpressure (Code = 89).	 Ensure use of correct cuff size, correct positioning and fit Ensure that the hose is not bent or pinched Ensure that the cuff and hose are not beneath patient After completing the above steps, attempt another measurement
90*	Unable to continue Brachial Assessment due to hardware problem (Code = 90).	 Turn the device OFF. Ensure the data/power input are connected Turn the device ON After completing the above steps, attempt another measurement
103*	File "DatabaseUpgradeReport.csv" is in use by other application (code=103)	Close the application
608*	Unable to backup database to <pathname> (Code = 608).</pathname>	• Select a different folder for backup (preferably at the local PC instead of the network), and then attempt to back up the database again
609*	Unable to restore database from <pathname> (Code = 609).</pathname>	 Copy the backup file in a different folder (preferably at the local PC instead of the network) Try to restore from the new location of the backup file.
611*	Unable to migrate data due to database error (Code = 611).	 Ensure that the PC can locate the SQL data server. Restart the SphygmoCor XCEL software.

Error Code	Error Message	Action
800*	Database server is not responding	Restart the computer
	(coue - 800).	Select the SphygmoCor XCEL installation media
		• Select Create Database and follow the instructions on the screen.
2046	Unable to continue Capture due to	• Ensure that the device is connected to the computer.
	2046).	• Ensure the cuff is connected to the device.
		 Ensure that there aren't any tubing obstructions
		• Turn the device OFF and ON.
		Attempt Capture again.
2047	Unable to continue Capture due to	Ensure that the device is connected to the computer
	2047). "Press the STOP button on	• Ensure that the cuff is connected to the device.
	the module to deflate the cuff	 Ensure use of the correct cuff size and that it is correctly positioned
		• Turn the device OFF and ON.
		Attempt Capture again.
2049	Unable to continue Capture due to	• Ensure the device is connected to the computer.
2050	2049).	• Ensure the cuff is connected to the device.
	(Code = 2050)	 Ensure use of the correct cuff size and that it is correctly positioned
		• Turn the module OFF and ON.
		Attempt Capture again.
2051	Unable to continue Capture due to	Press the STOP button on the device to deflate the cuff.
2053	2051) (Code = 2053)	• Ensure that the device is connected to the computer.
		• Ensure that the cuff is connected to the device.
		 Ensure use of the correct cuff size and that it is correctly positioned
		• Turn the device OFF and ON.
		Attempt Capture again
2058	Unable to continue Capture due to exceeding the maximum number of inflation (dofinition such a set	 Ensure that the patient's limb with cuff is stationary during measurement.
	measurement (Code = 2058).	• Ensure use of correct cuff size, correct positioning and fit.
		 Ensure that there are no obstructions such as thick clothing between the cuff and arm.
		• For PWV mode: Ensure the tonometer is detecting pulses.

Error Code	Error Message	Action		
		• After completing the above steps, attempt Capture again.		
2076	Unable to continue Capture due to	Press the STOP button on the device.		
	Module (Code = 2076).	• Ensure that the device is connected to the computer.		
		 In the Setup screen, click on the System and select Find Module. 		
		• Attempt Capture again.		
2077	Unable to continue Capture due to	Press the STOP button on the device.		
2078	(Code=2077)	• Turn the device OFF and ON.		
	(Code=2078)	• Ensure that the device is connected to the computer.		
		• Exit the XCEL software		
		• Start the XCEL software		
		 In the Setup screen, click on the System and select Find Module. 		
		Attempt Capture again.		
2079	Unable to continue Capture due to	• Ensure that the patient's limb with cuff is stationary.		
	denation of curr (code – 2079).	 Ensure use of the correct cuff size and that it is correctly positioned. 		
		 Ensure that there are no obstructions between the cuff and arm. 		
		• After completing the above steps, attempt Capture again.		
2080	Unable to continue Brachial Assessment due to brachial pressure being outside the normal range	 Ensure systolic pressure is within the range 50 to 260 mmHg and diastolic pressure is within the range 40 to 200 mmHg. 		
	(Code=2080)	• Ensure use of correct cuff size, correct positioning and fit.		
		 Ensure that there are no obstructions between the cuff and arm. 		
		• After completing the above steps, repeat measurement.		
2081*	Unable to continue Brachial	• Turn the module OFF/ON .		
	Assessment due to timeout.	Click OK button.		
		• Proceed with the measurement.		
2090*	One of these messages related to	• In the Set Up screen, click System Key.		
	appear:	• Enter the correct system key provided by AtCor Medical.		
	(a) Unable to find the Electronics Module due to mismatch serial number,			

Error Code	Error Message	Action
	 (b) Unable to start Brachial Assessment due to mismatch serial number (c) Unable to start Capture due to mismatch serial number 	
2091	 One of these messages related to "incomplete calibration" error will appear: (a) Unable to find the Electronics Module due to incomplete calibration (b) Unable to start Brachial Assessment due to incomplete calibration (c) Unable to start Capture due to incomplete calibration 	 Close SphygmoCor XCEL software. The SphygmoCor XCEL Electronics Module need to be calibrated. Please refer to XCEL Calibration KIT or contact AtCor Medical for technical support.
3051*	Unable to generate PWV Report (Code= 3051).	 Ensure that the entered distance in mm is correct. Ensure use of the correct cuff size and that it is correctly positioned. Ensure waveform quality (use the guidance bar). After completing the above steps, repeat PWV measurement.
3052*	Unable to generate PWV Report (Code= 3052).	 Ensure there is enough memory in the computer. Ensure the user has read/write permission for the computer on which the SphygmoCor XCEL software installed. Ensure the entered distance in mm is correct. Ensure use of the correct cuff size and that it is correctly positioned. Ensure waveform quality (use the guidance bar). After completing the above steps, repeat PWV measurement.
3053*	Unable to generate PWA Report (Code= 3053).	 Ensure use of correct cuff size, correct positioning and fit. Ensure waveform quality (use the guidance bar). After completing the above steps, repeat PWA measurement.
3054*	Unable to generate PWA Report (Code= 3054).	 Ensure there is enough memory in the computer. Ensure that the user has read/write permission where the XCEL software installed.

Error Code	Error Message	Action
		• Ensure use of correct cuff size, correct positioning and fit.
		 Ensure waveform quality (use the guidance bar).
		 After completing the above steps, repeat PWA measurement.
3060	Unable to start Capture (Code= 3060).	• Ensure that the device is connected to the computer.
		• Ensure that the cuff is connected to the device.
		• Ensure use of correct cuff size, correct positioning and fit.
		• Turn the device OFF and ON.
		• After completing the above steps, attempt Capture again.
3101*	Unable to complete the operation due to system error (Code=3101)	Close SphygmoCor XCEL software.
		• Turn the device OFF and ON.
		Open SphygmoCor XCEL software.
3125	Unable to calculate PWV (Code= 3125).	• Ensure that the entered distance in mm is correct.
3126*	Unable to find the Electronics Module (Code = 3126)	• Ensure the device is on.
	(0000 0120).	• Ensure the device is connected to the computer.
		• In the Setup screen, click System and select Find Module .
3128	Unable to continue due to unsupported screen resolution (Code	• Ensure the screen resolution is set to 1366 by 768 pixels or higher.
	=3128). It must be equal to or higher than 1366 by 768 pixels.	 You may need to adjust the window display scaling level to 100% via Control Panel → Appearance and Personalization → Display
3136*	Unable to continue due to the incompatible database version (code=3136)	Upgrade SphygmoCor XCEL software.

*If the problem persist persists, please contact AtCor Medical Technical support.

Maintenance

Database Manager

Database Manager allows you to backup and restore databases. You should backup the database regularly to protect against the possibility of losing important patient data. To access the Database Manager, click **Database** at the top of the screen.

System Da	atabase Help						- 20
Mode PWA	Backup Restore		Setup	• Capture Report			SphygmoCor XCEL
	Patients 5				Height: cm	Notes:	A
Patient ID	Last Name	First Name	Date Of Birth	Gender 📥	• <u> </u>		
1 Smith	ı	John	1/1/1954	Male			
2 Doe		Jane	4/7/1970	Female			Ψ.
3 Roe		Jane	10/16/1966	Female			
4 Smith	1	Johnnie	7/8/1998	Male	Brachial	Assessment	
Patient ID: 5 First Name: Jan Last Name*: Do	nie nie	Date of Birth*: Gender*	Dec 30 Female	Hide 1997	SYS: mmHg PP: mmHg		
New Searc	ch Edit Delete	e Save Cancel			Start		
						Sp Att	hygmoCor 🍲 Cor Medical

Backup a Database

1. Select Database - Backup.

🤯 Database backup		
💽 🍚 – 📜 🕨 Com	nputer > - 49 Search	n Computer 🔎
Organize 🔻		
 ★ Favorites ■ Desktop ▶ Downloads ™ Recent Places ■ Libraries ■ Documents ↓ Music ■ Pictures ■ Subversion ₩ Videos 	 Hard Disk Drives (3) Local Disk (C:) 288 GB free of 442 GB 288 GB free of 442 GB 1.95 GB free of 1.98 GB HP_RECOVERV (G:) 3.24 GB free of 21.0 GB Devices with Removable Storage (1) DVD RW Drive (F:) Network Location (7) 	
19 Computer 실 Local Disk (C:)	Clinical\$ (\\sfs) (H:)	-
File name: 📕 Save as type: 🏝	Atcor-xcell2.0 .bak	▼ ▼
) Hide Folders	Sa	ve Cancel

2. Choose a location to save your back up file and click **Save**.

The backup database must not be saved in any of the following locations:

- C:\
- C:\Program Files
- C:\Documents and Settings
- Desktop
- My Documents
- Any network folder

Restore a Database

The Database – Restore feature is used if a database becomes corrupted or needs to be installed onto another PC. The user can restore a backup database to the existing database. To delete the existing database, restore a blank database.

Note: For multi-user SphygmoCor XCEL, the software should not be accessed by other users before Restore is complete.

1. Select **Database - Restore**.



- 2. Click **Yes** if you are certain that you want to replace your existing database with a backup database.
- 3. Navigate to the location of the backup file and click **Open**. The restore will be completed automatically.

Note: To delete the existing database, go to C:\Atcor\XCEL 1.3\factory \rightarrow Database_BLANK.bak and restore existing database with this blank database.

Note: When restoring a SphygmoCor XCEL database from an earlier version, a message will appear indicating that the earlier version of the SphygmoCor XCEL database is backed up.

Basic System Care

Stability

Place the SphygmoCor XCEL device gently onto a stable surface. Dropping the unit can cause damage and result in the unit not operating correctly.

Choose a proper location that is free of clutter and away from high activity areas where the device could be accidentally dropped.

Pressure or impact damage

Do not apply heavy pressure to the SphygmoCor XCEL device or subject it to strong impact. Excessive pressure or impact can damage electronic components or otherwise cause malfunctions.

Temperature

The SphygmoCor XCEL System should be used and stored in ambient temperature and relative humidity within the range outlined in Technical Specifications.

Caution: Do not expose the unit to dirt, moisture or dust. Exposing the unit to dust or moisture could cause it to fail. Do not place the unit in direct sunlight – prolonged sun or heat exposure may result in overheating and damage to internal components. If you place the unit near a window, leave it inside your car, or take it outside in direct sun, it may overheat and damage internal components.

Magnetic fields

Magnets, television sets, radios, large electric motors or any other source of strong magnetic fields such as MRI devices will affect the operation or may cause it to fail.

Caution: Do not place any components near strong magnetic fields.

Liquids

Liquids on or inside any components of the SphygmoCor XCEL System can cause irreversible damage.

Caution: Do not spill liquids on any component.

Weight

Weight applied to the top of the SphygmoCor XCEL device may cause the unit enclosure to crack and other parts to be damaged.

Caution: Do not place any objects on top of the SphygmoCor XCEL device. Items on top of the SphygmoCor XCEL device may obscure or inadvertently activate the Stop Button.

Movement

Sudden jolts can cause damage to the unit or the tonometer if it is not stored properly.

Caution: Do not shake or drop the unit.

Shutting down

Do not power off the PC until the SphygmoCor XCEL software and Windows Operating System have been shut down. Wait 4 seconds after turning the computer OFF before turning it on again.

Caution: Exit the SphygmoCor XCEL software prior to shutting down the computer.

Equipment Calibration

Warning: The SphygmoCor XCEL System requires calibration annually. Failure to do so, may result in the device being out of specification. This may result in inaccurate measurements.

When calibration is due the follow message will appear.



Calibration can be completed by the customer by using the XCEL Calibration Kit (purchased from AtCor Medical). Alternatively, Calibration can be completed by AtCor Medical by returning your SphygmoCor XCEL System to an AtCor Medical representative. Please contact your local AtCor Medical representative for further information.

Note: If a message appears indicating incomplete calibration, please refer to the SphygmoCor XCEL Calibration Kit instruction manual for recalibration or contact AtCor Medical for technical support.

Sphygn	noCor XCEL 🛛 🔍
	Unable to find the Electronics Module.
•	Incomplete calibration (Code = 2091).
	Please calibrate the module before starting a new measurement.
	OK

For repairs, refer to qualified service personnel as instructed by AtCor Medical. The device does not contain any user-serviceable or reusable parts. Disassembly of the device by unauthorised personnel voids any warranty conditions.

Cleaning Instructions

To clean the SphygmoCor XCEL device, first unplug it from the computer and the power outlet. Using a damp cloth with mild detergent, gently wipe the equipment. Do not use other cleaning agents. Ensure excess liquids are wiped immediately from the equipment.

Caution: Do not spray cleaning agents or liquid directly on any components.

Notebook Batteries

Ensure when using the SphygmoCor XCEL System on notebook computers running on rechargeable batteries that the batteries are fully charged. Do not use the system on low battery power. If the notebook is abruptly shut down, the SphygmoCor database may be corrupted. Consult the notebook manufacturer's user documentation regarding the safety and maintenance of the notebook rechargeable battery.

Tonometer Care

The tip of the tonometer is a delicate and sensitive device, and can be easily damaged if dropped or misused. Follow the guidelines below to ensure tonometer lifetime is maintained.

When the tonometer is not in direct use with the patient, protect the tonometer by placing it in the storage holder in the tray or in the temporary storage holder on top of the SphygmoCor XCEL device.

Do not use this tonometer with any other instrumentation other than that supplied by AtCor Medical. Do not use a tonometer from any other supplier. The SphygmoCor XCEL tonometer is intended to be used in conjunction with the AtCor Medical SphygmoCor XCEL device only, which has a floating (isolated) grounding system.

Disinfection Instructions for Tonometer and Cuffs

The SphygmoCor XCEL product is considered a "non-critical" device. Therefore a low-level disinfection method has been provided to assist users to disinfect the tonometer and cuffs, which are the only patient contacting component of the SphygmoCor XCEL System (see below).

For tonometer disinfection, use a 70% Isopropyl Alcohol (IPA) impregnated wipe or cloth for low-level disinfection. Allow a contact time of at least 5 minutes.

For cuff cleaning, refer to the cleaning instructions accompanied in the cuff packaging.

Caution: Do not immerse the tonometer in any liquid as this will damage the tonometer electronics. Do not use coarse cloths for wiping the tonometer as this will damage the sensitivity of the transducer.

PWV Thigh Cuff Care

Cleaning Cuffs: Removable covers on straight and contoured cuffs make cleaning easy. The cuff covers are made of Nylon and Velcro. Remove the bladder then wash the cuff cover in a washing machine on gentle cycle or by hand using mild soap. Open the top of the cuff and line dry only. When completely dry, reinsert the bladder. If bladder is contaminated, wash it in soapy water and rinse well without getting any liquid in the bladder or tubing.

Disinfecting Cuffs: Spray or wipe entire surfaces of cuff with disinfectant until wet. Allow the cuff to remain visibly wet for a minimum of 10 minutes to insure complete disinfection. Wipe dry with clean cloth. The following disinfectants have been tested for compatibility with Hokanson cuffs. The manufacturers do not claim effective disinfection on porous surfaces; we have been unable to find any disinfectants that claim disinfection on porous surfaces. Hydrogen peroxide, hydrogen peroxide with silver (Sanosil[®]), hydrogen peroxide, peroxyacetic acid, silver (Steriplex^o"), silver with citric acid (PureGreen24Tm & SpectraSanTM 24), Octyl decyl dimethyl ammonium chloride with dioctyl dimethyl ammonium chloride with didecyl dimethyl ammonium chloride with dimethyl benzyl ammonium chloride (ProtexTM), 99% isopropyl alcohol, or T-SprayTM. Note: Some disinfectants may cause the cuff blue colour to bleed into cuff labelling. This does not affect the cuffs performance.

WARRANTY: Cuffs are guaranteed for one year from the date of purchase.

PWA Brachial Cuff Care

- Follow the application instructions for use to ensure the correct size cuff for the patient. Failure to do so will adversely affect the accuracy of the reading.
- Avoid contact with the cuff, other than that of the patient's limb, while measurement is in progress. Do not compress or pinch cuff tubing.
- Promptly remove cuff from patient when monitoring is not in progress
- During frequent or extended measurement, check cuff site and limb to ensure proper blood flow. Do not apply cuff where circulation may be compromised.

Cleaning instructions

The following cleaning methods have been applied 20 times to the cuff without any apparent negative effects.

The cuff may be sprayed with a mild disinfectant solution (e.g. Cidezyme[®] ENZOL[®], or 10% bleach solution), rinsed with distilled water and line dry. Ensure that no liquid enters tubing.

OR

To machine wash the cuff, remove the bladder and fully engage the hook and the loop. Machine wash warm with a mild detergent (50 -130° F or 1 - 54° C) and line dry.

Reference Information

Refer to our website <u>www.atcormedical.com</u> for a current list of SphygmoCor publications.



Electromagnetic Compatibility (EMC) Warnings & Declarations

- The SphygmoCor XCEL System needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in this manual
- Portable and mobile RF communications equipment could affect the SphygmoCor XCEL System
- Use of tonometers, power supplies, accessories, and cables other than those specified or supplied by AtCor Medical may affect EMC compliance of the SphygmoCor XCEL System.

Warning: The SphygmoCor XCEL System is intended for use by healthcare professionals only. The SphygmoCor XCEL System may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as reorienting or relocating the SphygmoCor XCEL System or shielding the location.

The ShygmoCor XCEL System has been tested, and complies with no deviations to the below immunity and emissions standards.

Electromagnetic Emissions			
Туре	Compliance to standard	Electromagnetic environment	
RF emissions (conducted and radiated)	CISPR11 Group 1 Class B	Clinical or Research	
Harmonic distortion	IEC 61000-3-2 Class A	Clinical or Research	
Voltage fluctuations and flicker	IEC 61000-3-3	Clinical or Research	

Electromagnetic Immunity			
Туре	Compliance to standard	Test levels	
		Clinical or Research environment	
Electrostatic Discharge	IEC 61000-4-2	±8kV contact ±2kV, ±4kV, ±8kV, ±15kV air	
Radiated RF	IEC 61000-4-3	3V/m 80MHz-2.7GHz 80% AM @ 1kHz	
Proximity Fields	IEC 61000-4-3	385MHz, 18Hz pulse, 27V/m, 3m distance 450MHz, 18Hz pulse, 28V/m, 3m distance 710MHz, 217Hz pulse, 9V/m, 3m distance 745MHz, 217Hz pulse, 9V/m, 3m distance 80MHz, 217Hz pulse, 9V/m, 3m distance 810MHz, 18Hz pulse, 28V/m, 3m distance 930MHz, 18Hz pulse, 28V/m, 3m distance 1720MHz, 18Hz pulse, 28V/m, 3m distance 1720MHz, 217Hz pulse, 28V/m, 3m distance 1845MHz, 217Hz pulse, 28V/m, 3m distance 1970MHz, 217Hz pulse, 28V/m, 3m distance 2450MHz, 217Hz pulse, 28V/m, 3m distance 5240MHz, 217Hz pulse, 9V/m, 3m distance 5500MHz, 217Hz pulse, 9V/m, 3m distance 5785MHz, 217Hz pulse, 9V/m, 3m distance	
Electrical fast transients/bursts (EFT)	IEC 61000-4-4	±2kV 100kHz repetition frequency	
Surges	IEC 61000-4-5	Line to line ±0.5kV, ±1kV Line to earth ±0.5kV, ±1kV, ±2kV	

Conducted disturbances	IEC 61000-4-6	3Vrms, 150kHz to 80MHz 6Vrms for ISM bands between 150kHz-80MHz at 80%AM at 1KHz modulation
Power Frequency Magnetic Fields	IEC61000-4-8	30A/m @ 50Hz and 60Hz
Voltage Interruptions	IEC61000-4-11	0%, 0°, 5 seconds @ 100V 0%, 0°, 5 seconds @ 240V
Voltage Dips	IEC61000-4-11	70%, 0°, 0.5 seconds @ 100V 0%, 0°, 1 cycle @ 100V 0%, 45°, 90°, 135°, 225°, 270°, 315°, @ ½ cycle @ 100V 70%, 0°, 0.5 seconds @ 240V 0%, 0°, 1 cycle @ 240V 0%, 45°, 90°, 135°, 225°, 270°, 315°, @ ½ cycle @ 240V

Important information required by the FCC

Note: This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation. This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to radio or television reception, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

-Reorient or relocate the receiving antenna.

-Increase the separation between the equipment and receiver.

-Connect the equipment into an outlet on a circuit different from that to which the receiver is connected.

-Consult the dealer or an experienced radio/TV technician for help.

Technical Specifications

Physical and Environmental Specifications

Operating Ambient temperature:	+15°C to 40°C (59°F to 104°F)		
Operating Relative humidity:	15% to 95% non-condensing		
Storage Ambient temperature:	-20°C to 65°C (-4°F to 149°F)		
Transport Ambient temperature:	-20°C to 65°C (-4°F to 149°F)		
Storage Relative humidity:	20% to 90% non-condensing		
Transport Relative humidity:	20% to 90% non-condensing		
	Input	100-240 VAC, 50-60Hz	
Extornal Dowor supply	Output	15VDC at 2A	
External Power supply	Protective Class	IEC Class II, Double Insulated	
	Power Connector	4 PIN DIN	
	Enclosure Material	Polycarbonate	
	Weight	0.7 kg (1.5 lbs)	
Physical Specifications	Dimensions (EM4C)	9.9 (l) x 19 (w) x 17.2 (h) cm	
	Dimensions (Tray)	30.5(l) x 16.7 (w) x 8.6 (h) cm	
Measurement	NIBP	Oscillometric. Diastolic values correspond to Phase 5 Korotkoff sounds.	
	NIBP, PWA	Sys: 50 - 260 mmHg Dia: 40 – 200 mmHg	
Range	PWV	2 to 25 m/s	
	Heart rate	30 - 220 beats per minute	
	Display	0 - 300mmHg	
Resolution	1 mmHg		
Tonometer	Uncalibrated pressure transducer		

Maximum Intended Design Life

The Maximum Intended Design Life of the SphygmoCor XCEL device is 5 Years.

The Maximum Intended Design Life of the Tonometer is 3 Years.



Туре	IBM Compatible PC
Processor	Intel or compatible, Core i3, 32 or 64 bits
Nominal Speed	2.4 GHz minimum
Memory	2GB RAM minimum
Hard Disk	Size 40 GB
	2GB for Installation
	10GB for database
Accessories	DVD drive, USB port
Printer Drivers	User Provided
Communications	USB port
Minimum Display Resolution	1366 x 768 pixels
Operating Systems	Windows 7 Professional + SP1, or Windows 8 Pro, or Windows 10 Professional

Recommended Minimum Computer Requirements

Classification of SphygmoCor System

The SphygmoCor XCEL System is classified as follows:

- Class II ٠
- Double insulated •
- Type BF Equipment •
- This equipment is not suitable for use in the presence of a flammable anaesthetic mixture with air or ٠ with oxygen or nitrous oxide.
- **Continuous Equipment** •

The following symbols, listed below with their meanings, are used throughout this manual as well as on the equipment:



Consult Operating Instructions

Type BF Applied Part

Date of Manufacture





Standards

The SphygmoCor XCEL System is designed, tested and approved to the following safety and performance standards:

- IEC/EN 60601-1 ANSI/AAMI ES60601-1 CSA C22.2 No 60601.1 Medical electrical equipment -- Part 1: General requirements for safety.
- IEC 80601-2-30 Medical electrical equipment -- Part 2-30: Particular requirements for basic safety and essential performance of automated non-invasive sphygmomanometers.
- IEC 60601-1-2 Medical electrical equipment -- Part 1: General requirements for safety Section 2. Collateral Standard: Electromagnetic compatibility Requirements and tests.
- 47 CFR Part 15 Radio Frequency Devices, Subpart B (Unintentional Radiators)

SphygmoCor XCEL Electronics Module, model number EM4X

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

Changes or modifications not expressly approved by Atcor Medical Pty. Ltd., would void the user's authority to operate the product.



Explanation of Parameters/Indices

The SphygmoCor PWA report screens and printed reports contain key patient details, the averaged (derived) aortic waveform, and central hemodynamic parameters.

The following parameters apply to each averaged waveform:

Displayed Parameter Summary:

Calculated Parameters	Description	Abbreviation	Units	
Peripheral Pulse Indices				
Heart Rate	Number of heart beats per minute	HR	Beats/min	
Period	Cardiac pulse length in time	T⊧	milliseconds	
Ejection Duration	The ejection time in one pulse	ED	milliseconds	
Brachial Systolic and Diastolic Pressure	The highest and lowest peripheral (brachial) pressure respectively in one cardiac pulse	SYS, DIA	mmHg	
Pulse Pressure - Brachial	The difference between maximum and minimum peripheral pressure in one cardiac pulse. Pulse pressure represents the height of the pressure pulse.	РРР	mmHg	
Central Aortic Indices				
Aortic Systolic and Diastolic Pressure	The highest and lowest central aortic pressure respectively in one cardiac pulse	SP, DP	mmHg	
Aortic Pulse Pressure	The difference between maximum and minimum (SP, DP) central aortic pressure in one cardiac pulse. Pulse pressure represents the height of the pressure pulse.	PP	mmHg	
Mean Pressure	Average pressure in a cardiac pulse	MP	mmHg	
Augmentation Index at 75 BPM heart rate	Alx normalised to heart rate of 75 BPM	Alx@75	%	



Features of the Arterial Pulse



The following diagram shows the central aortic pressure waveform.

Parameter	Description	Symbol	Units	Calculation	Sample from above graph
Systolic Pressure	The maximum pressure	SP	mmHg	Maximum Point on Chart	150 mmHg
Diastolic Pressure	The minimum Pressure	DP	mmHg	Minimum Point on Chart	100 mmHg
Pulse Pressure	The height of the pressure pulse	РР	mmHg	PP = SP – DP	= 150 – 100 mmHg = 50 mmHg
Augmentation Pressure	The difference in pressure between the two peaks during heart ejection (systolic). The first peak (P_1) is related to cardiac ejection and the second peak (P_2) is related to wave reflection due to arterial stiffness.	AP	mmHg	$AP = P_2 - P_1$	= 150-140 mmHg = 10 mmHg
Augmentation Index	The difference between the two peaks during heart ejection, relative to the pressure pulse height or pulse pressure (PP). Alx is defined as (AP/PP) x 100%. The measurement is an indication of arterial wave reflection which is due to arterial system elasticity.	Alx	%	Alx=(AP/PP)x100%	= (10/50)*100% = 20 %

Warranty

The SphygmoCor[®] XCEL system, excluding the tonometer, is supplied with a full parts and labour warranty. (The length of this warranty varies based on statutory requirements in each country – contact your distributor to determine the warranty for your device). Any parts that fail during this warranty period will be repaired or replaced without charge, at the discretion of AtCor Medical. This excludes any damage to the instrument arising from operational wear and tear, or from misuse.

The tonometer has a 120-day parts and labour warranty.

Service and technical support for the SphygmoCor XCEL system will be provided by AtCor Medical technical support group. Contact your local distributor to organise the service.

Product Support

If you have additional questions after reviewing the manual, contact your local AtCor Medical representative / distributor. The details of your local representative / distributor can be found at www.atcormedical.com .

Please quote your *System Serial Number* to your Product Support Representative. This serial number can be found by opening the SphygmoCor XCEL software and selecting **Help>About** from the Main Menu. A window will appear and will contain the System ID Number. This window also contains the Version of software and the installed options.

When contacting your local AtCor Medical representative / distributor, please have your SphygmoCor XCEL System nearby as well as copy of the Operator's Manual open on your PC. You should be prepared to give the following information, if applicable:

- System ID number
- The error number and exact wording of any messages that appear. The error number will appear in the corner of the error message window.
- Relevant details related to the issue, and activity that transpired prior to the problem occurrence.
- Any action that you took as a result of a warning/error/problem.
- The version of your SphygmoCor software.

Safety Warnings

Medical Equipment Systems Compliance Warnings

(IEC60601-1 Section 16 & IEC60601-1-1)

IEC60601-1-1 compliance is the responsibility of the end user. To ensure compliance to IEC60601-1 Section 16 & IEC60601-1-1, the SphygmoCor XCEL System must meet the following three conditions:

- 1. Ensure that only the SphygmoCor XCEL device, Tonometer, cuffs and hose are within reach of the patient. The computer and other devices should be at least 1.5m (6ft) away from the patient.
- 2. The PC and peripherals (e.g. USB hubs) must comply with IEC60950 or equivalent.



- 3. The enclosure leakage current from any device within the patient environment, including any parts of equipment which extend into that environment, is not more than 0.1mA in normal condition and 0.5mA in the single fault condition of interrupting an earth conductor in any single power supply cord (for the U.S.A. the single fault limit is reduced to 0.3mA).
- 4. The required low enclosure leakage current may be achieved by powering the PC and peripherals from an isolation transformer. Connecting the SphygmoCor XCEL System to other non-isolated monitoring equipment or communication networks is not recommended. It is the end user's responsibility to ensure compliance with IEC60601-1 Section 16 and IEC60601-1-1.

General Safety Warnings

- 1. When using the SphygmoCor XCEL device, do not connect the power cable of the computer to multiple portable socket-outlets or power boards which are connected to other devices. Do not place the multiple portable socket-outlet or power board on the floor while the SphygmoCor XCEL System is in use.
- 2. Do not connect any peripheral devices (e.g. printer, externally powered USB hubs) to the computer while using the SphygmoCor XCEL System as they may breach the patient isolation requirements of IEC60601-1 & IEC60601-1-1.
- 3. When using an isolation transformer and a multiple portable socket-outlet or power board to connect to the computer and the SphygmoCor XCEL System, do not connect any other devices or equipment to the multiple portable socket-outlet or power board.
- 4. When using the SphygmoCor XCEL System, the operator should not touch the computer and the patient at the same time.
- 5. Do not connect or use any cables, cuffs, hose or tonometer other than those specified for use with the SphygmoCor XCEL System.
- 6. Do not disassemble the SphygmoCor XCEL device. The SphygmoCor XCEL device does not contain any serviceable parts. Servicing shall be performed by qualified service personnel.

USA Privacy Rule

This AtCor Medical product stores, displays and exports patient health information which could affect HIPAA compliance. It is the responsibility of the health care organization that is subject to the Privacy Rule to ensure compliance with HIPAA regulations. AtCor Medical does not make any claim with respect to compliance with HIPAA for its products. The Privacy Rule (also known as Standards for Privacy of Individually Identifiable Health Information) is contained in Title 45 of the Code of Federal Regulations, Part 160 and Subparts A and E of Part 164.

Cybersecurity

If the PC used with the SphygmoCor XCEL System is connected to a network or the internet, the provider of services is responsible for ensuring the security of the information.

Disposal

According to the Official Journal of the European Union WEEE Directive 2002/96/EC that requires the proper disposal of electrical and electronic equipment. This device should be disposed of separately, not as unsorted municipal waste. To dispose of your device, you should use appropriate collection, reuse and recycling systems available in your region. The use of these collection, reuse and recycling systems is

designed to reduce pressure on natural resources and prevent hazardous substances from damaging the environment.

If you need information on these disposal systems, please contact your local waste administration. The crossed-bin symbol invites you to use these disposal systems. If you require information on collection and disposal of your AtCor Medical device please contact AtCor Medical Head Office or local distributor.




Appendix

Installing SphygmoCor XCEL in a Networked Configuration

By default, SphygmoCor XCEL is set up to operate on a single PC where the SQL Database and the SphygmoCor XCEL Software are operating on the same computer.

Users who require the SQL Server Database to be accessed by multiple computers on the same local network can follow the guide below to configure the SphygmoCor XCEL system to function in a networked environment.



Features Installed on each Client:

SphygmoCor XCEL

Warning: Depending on your database size and network configuration, the database Backup and Restore functions described in section *Database Manager* may not be successful over a network connection. This could lead to database corruption. It is strongly recommended that the Backup and Restore functions **not** be used from the XCEL client over a network. These database operations should be carried out on the Server computer by an experienced SQL Server user via SQL Server Management Studio. It is strongly recommended to regularly backup the XCEL database.

Note: Network Installation steps should be implemented by an IT administrator.

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Preparation

- 1. Administrative access to the Server, or host computer.
- 2. Administrative access to each Client networked computer.
- 3. Ensure that Microsoft SQL 2014 Server is not installed on the Client.

Installing Microsoft SQL Server and AtCor Database on the Server

- 1. Select the SphygmoCor XCEL installation media in the Server computer and launch 'setup.exe'.
- 2. Go through the installation steps and when asked to select a Setup Type, select Custom
- In the Select Features dialog, only select the feature "Microsoft SQL Server and AtCor Database" for installation. Deselect all other features. Refer to the *Custom Installation* section for more details.
- 4. Continue to complete installing Microsoft SQL Server and AtCor database.

Configuration of the Server

Note: The configuration settings below may vary depending on your computer and network configuration.

Follow the steps below after installing SQL Server and the AtCor Database on the Server computer:

- a. From the Windows Start menu go to All Programs > Microsoft SQL Server
 2014 > Configuration Tools > SQL Server 2014 Configuration Manager.
- b. In the left pane expand and select SQL Server Configuration Manager > SQL Server Network Configuration > Protocols for ATCORSERVER2014.
- c. In the right pane, set **TCP/IP** to enabled if it is not, then right-click on **TCP/IP** and select **Properties.**
- d. In the **TCP/IP Properties** dialog, go to the **IP Addresses** tab and scroll down to the **IPAII** group.
- e. In the TCP Port box, enter an available port number (for example 49394). If there is a dynamic port assigned in the TCP Dynamic Ports box, you may use this value for the TCP Port. Make a note of the port number you assigned as it will be used in the steps below. Port numbers can be assigned between 49152 to 65535.
- f. Delete any value in the **TCP Dynamic Ports** box. (the box must be empty, do not use a value of 0).

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g. Click **Apply** and then **Ok** to apply the change.

- h. Click **OK** to the warning that the service must be restarted before the changes take effect.
- i. In the left pane, select SQL Server Services.
- j. In the right pane, right-click SQL Server (ATCORSERVER2014) and select Restart.
- k. Close the SQL Server Configuration Manager.
- I. On the **Start** menu, click **Run**, type **WF.msc** in the run window.
- m. Click OK. This should open Windows Firewall with Advanced Security.
- n. In the left pane, right-click **Inbound Rules**, and then click **New Rule**.
- o. In the Rule Type dialog, select Port and click Next.
- p. In the **Protocol and Ports** dialog box, select **TCP**. Select **Specific local ports** and type in the port number noted in step e, then click **Next**.
- q. Select Allow the connection if it is secure, and then click Next.
- r. Select any profiles that would meet to your network security needs when accessing the AtCor Database Engine then click **Next**.
- s. In the **Name** dialog box, type a name (e.g. "AtCor DB TCP port") and optional description for this rule, then click **Finish**.
- Create another inbound port rule for the UDP port 1434. In the Protocol and Ports dialog box, select UDP. Select Specific local ports and enter 1434 as the port number then click Next. Repeat steps q to s (In the Name dialog box, type a name e.g "AtCor DB UDP port").
- u. Close Windows Firewall with Advanced Security.
- v. Restart the computer.

Installing SphygmoCor XCEL on each Client

- 1. Select the SphygmoCor XCEL installation media in the Client computer and launch 'setup.exe'.
- 2. Go through the installation steps and when asked to select a Setup Type, select Custom.
- 3. In the Select Features dialog, only select the feature "**SphygmoCor XCEL**" for installation. Deselect all other features. Refer to the *Custom Installation* section for more details.
- 4. Continue to complete installing SphygmoCor XCEL.

Connecting to network database using SphygmoCor XCEL

- 1. Launch the SphygmoCor XCEL software.
- From the drop-down box select the applicable Server name (Host computer's name) and press OK.

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- 3. SphygmoCor XCEL will now be connected to the selected Networked Database.
- 4. Repeat this process for each Client computer.

Installing Recommended Fonts on Different Versions of Windows

The following error message (Code=3251) indicates the unavailability of one or more fonts in Windows. In this case, for example, the **recommended** font (MS PGothic) is not available and the user needs to add this font to Windows.



To resolve this error, follow the instructions below depending on the operating system used.

Windows 7 and Windows 8

- 1. Go to the following website to download the recommended font.
- https://www.microsoft.com/typography/fonts/product.aspx?PID=161
- 2. Right click on the downloaded font and click Install.

Windows 10

- 1. Click the **Start** button.
- 2. In Settings, click System.
- 3. Click Apps & features.
- 4. Click on the link, Manage optional features.
- 5. Click on the "+" icon next to Add a feature.
- 6. Scroll to find your recommended font. Click on that item, then click on Install.
- 7. Click on the back arrow in the upper left corner of the window to see the font installation in progress.
- 8. When finished, click **Set optional feature history** to confirm installation was successful.
- 9. Restart Windows.

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