

## EN English Introduction

Blood pressure measurements determined with X1 are equivalent to those obtained by a trained operator using cuff/stethoscope auscultation method when the limits prescribed by the American National Standard, Electronic or Automated Sphygmomanometry. This unit is to be used by adult consumers in a home environment. The patient is an intended operator. Do not use this device on infants or neonates. X1 is protected against manufacturing defects by an established International Warranty Program. For information you can contact the manufacturer, Rossmax International Ltd. Attention: Consult the accompanying documents. Please read this manual carefully before use. For specific information on your own blood pressure, contact your physician. Please be sure to keep this manual.

### Real Tuffing Measuring Technology

This unit uses the oscillometric method to detect your blood pressure. Before the cuff starts inflating, the device will establish a baseline cuff pressure equivalent to the air pressure. This unit will determine the appropriate inflation level based on pressure oscillations, followed by cuff deflation.

During the deflation, the device will detect the amplitude and slope of the pressure oscillations as they determine for you the systolic blood pressure, diastolic blood pressure, and pulse.

### Preliminary Remarks

This Blood Pressure Monitor complies with the European regulations and bears the CE mark, "CE 1639". The quality of the device has been verified and conforms to the provisions of the technical directive 93/42/EEC (Medical Device Directive), Annex I essential requirements and applied harmonized standards.

EN 1060-1: 1995/A2: 2009 Non-invasive sphygmomanometers - Part 1 - General requirements. EN 1060-3: 1997/A2: 2009 Non-invasive sphygmomanometers - Part 3 - Supplementary requirements for automatic blood pressure measurement systems. EN 1060-4: 2004 Non-invasive sphygmomanometers - Part 4 - Test Procedures to determine the overall system accuracy of automated non-invasive sphygmomanometers. ISO 81060-2:2013 Non-invasive sphygmomanometers - Part 2: Clinical investigation of automated measurement systems.

The blood pressure monitor is designed for long service time. Ensure continued accuracy. It's recommended that digital blood pressure monitors require recalibration. This monitor (under normal usage with approx. 3 measurements a day) does not require recalibration for 2 years. Once the unit should be re-calibrated the device will display **CR**. The unit should also be re-calibrated if the monitor sustains damage due to burst or bubble formation, or extreme heat or cold temperatures/humidity changes. When **CR** appears, simply return to your nearest dealer for recalibration service.

### Blood Pressure Standard

Refer to the definitions of the World Health Organization, the blood pressure ranges can be classified as follows: (Ref. 1993 WHO Hypertension Guidelines for the management of hypertension). This blood pressure classification are based on historical data, and may not be directly applicable to any particular patient. It is important that you consult with your physician regularly. Your physician will tell you your normal blood pressure range as well as the point at which you will be considered at risk. For reliable monitoring and recording of blood pressure, keeping long term records is recommended. Please download the blood pressure log at our website www.rossmax.com.

### Display Explanations

**EE / Measurement Error:** Make sure the L-plug is securely connected to the air socket and the cuff correctly and keep arm steady during measurement. If the error keeps occurring, return the device to your local distributor or service center.

**E1 / Air Circuit Abnormality:** Make sure the L-plug is securely connected to the air socket on the side of the unit and measure again quietly. If the errors still occur, return the device to your local distributor or service center for help.

**E2 / Pressure Exceeding 300 mmHg:** Switch the unit on the error measure again quietly, if the error keeps occurring, return the device to your local distributor or service center.

**E3 / Data Error:** Remove the batteries, wait for 60 seconds, and reload. If the error keeps occurring, return the device to your local distributor or service center.

**Er / Exceeding Measurement Range:** Measure again quietly, if the error keeps occurring, return the device to your local distributor or service center.

### Cuff Wrap Detection

If the cuff was wrapped too loosely, it may cause unreliable measurement results. The cuff can help to determine if the cuff is wrapped snugly enough. The specified **CO** appears on the display if "loosen cuff" has been detected during measurement. Otherwise the specified **CO** appears if the cuff is wrapped correctly during measurement.

### Movement Detection

The "Movement Detection" helps reminding the user to remain still and is indicating any movement during measurement. The specified icon appears once a "body movement" has been detected during and after each measurement.

Note: It's highly recommended that you measure again if the **CO** appears.

### Hypertension Risk Indication (HR)

The World Health Organization, classifying blood pressure ranges into 6 grades. This unit is equipped with innovative blood pressure risk indication, which visually indicates the actual risk level (optimal / normal / high-normal / grade 1 hypertension / grade 2 hypertension / grade 3 hypertension) of the result after each measurement.

**Irregular Heartbeat (HRB) Detection** This unit is equipped with an Irregular Heartbeat (HRB) Detection which allows those who have an irregular heartbeat to obtain accurate measurements alerting the user of the irregular heartbeat.

Note: It is strongly recommended that you consult your physician if the HRB icon **HR** appears often.

### Using the AC Adaptor (Optional)

1. Connect the AC adaptor with the AC adaptor jack on the right side of the unit. 2. Plug the AC adaptor into the power outlet. (AC adaptors with required voltage and current indicated on the AC adaptor jack).

⚠ Please unplug the batteries when operating with the AC mode for a longer period of time. Leaving the batteries in the compartment for a long time may cause them to overheat and possibly catch fire.

2. Batteries are not needed when operating with the AC mode. 3. AC adaptors are optional. Please contact the distributor for the compatible AC adaptors. 4. Use only the authorized AC Adaptor for this APPENDIX 1.

### Installing Batteries

1. Press down and lift the battery cover in the direction of the arrow to open the battery compartment. 2. Install or replace 4 "AAA" sized batteries in the battery compartment according to the directions indicated on the battery cover. 3. Replace the battery cover by clicking in the bottom hooks first, then push in the top end of the battery cover.

⚠ Do not service or maintain device and cuff while in use. 2. Do not use the tubing and/or AC adaptor for any other purpose than those specified, as they can cause risk of strangulation. 3. Do not use service or maintain device and cuff while in use.

**Measurement Method** Oscillometric. Pressure: 30-260 mmHg; Pulse: 40-199 beats/minute

**Pressure Sensor** Semi-conductor. Accuracy: ±3 mmHg; Pulse: ±5% of reading

**Inflation Pump Drive** Automatic Air Release Valve. Memory capacity: Last Number Memory Recall. Auto-shut-off: 1 minute after last key operation

**Permissible Operating Temp** 10°C-40°C (50°F-104°F); 15%-85% RH. **Relative and Humidity** 15%-85% RH. **Permissible Transport and Store** -10°C-60°C (-14°F-140°F); 10%-90% RH; **air temperature and Humidity** 700-1060 hPa

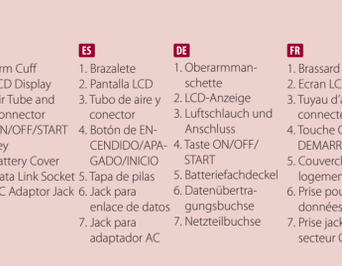
**Power Source** DC 6V four AAA Batteries. **AC Power Source** Plug type: outer(-) is 04.0, inner(+ ) is 01.7.

**Dimensions** 85 (L) X 1298 (W) X 682 (H) mm. **Weight** 226.0g (G.W) (w/o Batteries).

**Arm circumference** Adult: 24-40 cm (9.4"-15.7"). **Type** BF: Device and cuff are designed to provide special protection against electrical shocks.

**IP Classification** IP21: Protection against harmful ingress of water. **IP21**: Protection against harmful ingress of water.

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## ES Español

Las mediciones de presión arterial determinadas con X1 son equivalentes con aquellas obtenidas por un observador entrenado usando el método de auscultación con brazalete/estetoscopio dentro de los límites especificados en el norma ANSI para esfigmomanómetros electrónicos o automatizados. Esta unidad se ha concebido para ser usada por adultos en un entorno doméstico. El paciente es el operador previsto. No use este aparato para niños o recién nacidos. El está protegido contra defectos de fabricación por un programa de garantía internacional. Para una información acerca de la garantía, usted puede contactar al fabricante, Rossmax International Ltd. Atención: Consulte los documentos acompañados. Por favor, lea este manual cuidadosamente antes de usarlo. Para información específica acerca de su propia presión arterial, contacte a su médico. Por favor, fíjese en guardar este manual.

**Tecnología de medición real tuffing** Esta unidad usa el método oscilométrico para detectar su presión arterial. Antes de que el brazalete comience con el inflado, el aparato establece la presión de referencia del brazalete equivalente a la presión atmosférica. Esta unidad determinará el nivel de flado apropiado basándose en oscilaciones de la presión seguida por el desinflado del brazalete. Durante el desinflado, el aparato detectará la amplitud y la pendiente de las oscilaciones de presión que determinan para usted la presión sistólica, la presión arterial diastólica y el pulso.

**Notas preliminares** El monitor de presión arterial está conforme con las disposiciones europeas y lleva la marca "CE 1639". La calidad del aparato ha sido verificada y está conforme con la Directiva europea 93/42/CEE (Directiva de Aparatos Médicos), Anexo I requisitos esenciales y normas armonizadas aplicadas.

EN 1060-1: 1995/A2: 2009 Esfigmomanómetros no invasivos - Parte 1 - Requisitos generales. EN 1060-3: 1997/A2: 2009 Esfigmomanómetros no invasivos - Parte 3 - Requisitos suplementarios aplicables a los sistemas electromecánicos de medición de la presión sanguínea.

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ISO 81060-2:2013 esfigmomanómetros no invasivos - Parte 2: Investigación clínica sobre dispositivos de medición de la presión sanguínea.

Este monitor de presión arterial ha sido diseñado para una larga vida útil. Para garantizar una exactitud permanente, se recomienda que todos los montores digitales de presión arterial requieran recalibraciones. Este monitor (con un uso normal de aprox. 3 mediciones al día) no requiere recalibración por 2 años. Una vez que el dispositivo debe ser recalibrado se mostrará el código de error **CR**. La unidad también debería ser recalibrada si el monitor sustains daño debido a formación de burbujas o a temperaturas extremadamente calientes o frías / cambios de humedad.

Nota: Si el error se repite, simplemente devuélvalo al distribuidor más cercano para el servicio de recalibración.

**Troubleshooting** If any abnormality will arise during use, please check the following points.

**Symptoms** Check points Correction. No display when the batteries run down. Replace them with four new batteries. Have the batteries/ polarities been positioned incorrectly? Re-insert the batteries in the correct positions. Wrap the cuff properly so that it is positioned correctly. Did you talk or move during measurement? Measure again. Keep arm steady during measurement. Did you vigorously shake the cuff during measurement?

Note: If the unit still does not work, return it to your dealer. Under no circumstance should you disassemble and repair the unit yourself.

**Cautionary Notes** 1. The unit contains high-precision assemblies. Therefore, avoid extreme temperatures, humidity, and direct sunlight. Avoid dropping or strongly shaking the main unit, and protect it from dust. 2. Do not touch the blood pressure monitor body and the cuff carefully with a slightly damp, soft cloth. Do not press. Do not wash the cuff or use chemical cleaner on it. Never use ether, alcohol or petrol (gasoline) as cleaner.

3. Leaky batteries can damage the unit. Remove the batteries when the unit is not used for a long time. 4. The unit should not be freed by children to avoid hazardous situations. 5. If the unit is stored near freezing, allow it to acclimate at room temperature before use. 6. This unit is not field serviceable. You should not use any tool to open the device nor should you attempt to adjust anything inside the device. If you have any problems, contact your local distributor or service center for help.

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**Using the AC Adaptor (Optional)** 1. Connect the AC adaptor with the AC adaptor jack on the right side of the unit. 2. Plug the AC adaptor into the power outlet. (AC adaptors with required voltage and current indicated on the AC adaptor jack).

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**Measurement Method** Oscillometric. Pressure: 30-260 mmHg; Pulse: 40-199 beats/minute

**Pressure Sensor** Semi-conductor. Accuracy: ±3 mmHg; Pulse: ±5% of reading

**Inflation Pump Drive** Automatic Air Release Valve. Memory capacity: Last Number Memory Recall. Auto-shut-off: 1 minute after last key operation

**Permissible Operating Temp** 10°C-40°C (50°F-104°F); 15%-85% RH. **Relative and Humidity** 15%-85% RH. **Permissible Transport and Store** -10°C-60°C (-14°F-140°F); 10%-90% RH; **air temperature and Humidity** 700-1060 hPa

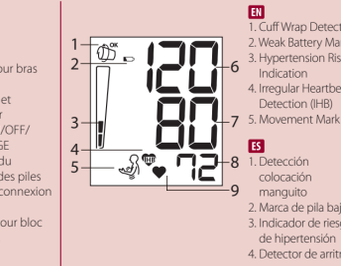
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## DE Deutsch

Die Messungen der Blutdruckwerte mit dem X1 sind äquivalent zu den durch einen geschulten Beobachter mit dem auskultatorischen Blutdruckmessgerät mit Manschette und Stethoskop gemessenen. Dieses Gerät ist für erwachsene Verbraucher in häuslicher Umgebung vorgesehen. Der Patient ist für die Anwendung des Gerätes nicht in Frage kommen oder kleinkindern anzuwenden. Das Modell X1 ist durch ein etabliertes internationales Garantieprogramm gegen Herstellerfehler geschützt. Für Garantieinformationen wenden Sie sich bitte an den Hersteller. Bitte lesen Sie die Bedienungsanleitung sorgfältig durch. Spezifische Informationen zu Ihrem eigenen Blutdruck erhalten Sie von Ihrem Arzt. Bitte bewahren Sie diese Bedienanleitung auf.

**Real-Fuzzy-Messungstechnik** Diese Technologie Ihres Blutdruckes die oszillometrische Methode. Bevor die Manschette aufgepumpt wird, ermittelt das Gerät ein Manschettenäquivalent für den Luftdruck. Das Gerät entscheidet anhand der Druckschwankungen über den Aufpumpdruck und führt danach eine Druckentlastung der Manschette durch. Während der Druckentlastung erkennt das Gerät die Amplitude und Flanken der Druckschwankung und ermittelt daraus für Sie den systolischen Blutdruck, den diastolischen Blutdruck und den Puls.

**Vorläufige Anmerkungen** Diese Vorläufige Anmerkungen erfüllen die europäischen Vorschriften und trägt das CE-Kennzeichen "CE 1639". Die Qualität des Geräts wurde überprüft und entspricht den Forderungen der Richtlinie des EU-Rates 93/42/EEC (Medizinärger Richtlinie). A) wesentliche Anforderungen, sowie den entsprechenden harmonisierten Normen.

EN 1060-1: 1995/A2: 2009 Nichtinvasive Blutdruckmessgeräte - Teil 1 - Allgemeine Anforderungen. EN 1060-3: 1997/A2: 2009 Nichtinvasive Blutdruckmessgeräte - Teil 3 - Zusätzliche Anforderungen an die Genauigkeit der Messung.

EN 1060-4: 2004 Nichtinvasive Blutdruckmessgeräte - Teil 4: Testprozeduren zur Bestimmung der Gesamtmesstauferlässlichkeit automatischer, nichtinvasiver Blutdruckmessgeräte (ISO 81060-2:2013 Nicht-invasive Sphygmomanometer - Teil 2: Klinische Untersuchung der automatisierten Blutdruckmessgeräte).

ISO 81060-2:2013 esfigmomanómetros no invasivos - Parte 2: Investigación clínica sobre dispositivos de medición de la presión sanguínea.

Este monitor de presión arterial ha sido diseñado para una larga vida útil. Para garantizar una exactitud permanente, se recomienda que todos los montores digitales de presión arterial requieran recalibraciones. Este monitor (con un uso normal de aprox. 3 mediciones al día) no requiere recalibración por 2 años. Una vez que el dispositivo debe ser recalibrado se mostrará el código de error **CR**. La unidad también debería ser recalibrada si el monitor sustains daño debido a formación de burbujas o a temperaturas extremadamente calientes o frías / cambios de humedad.

Nota: Si el error se repite, simplemente devuélvalo al distribuidor más cercano para el servicio de recalibración.

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## FR Française

Les mesures de pression artérielle réalisées avec l'X1 sont équivalentes à celles obtenues par un observateur expérimenté utilisant un brassard/stéthoscope, dans les limites prescrites par la norme nationale américaine sur les sphygmomanomètres électroniques ou automatiques. Cet instrument est réservé à un usage domestique par des adultes. Le patient est un opérateur prévu. Ne l'utilisez pas pour évaluer la pression artérielle d'enfants ou de bébés. X1 est garanti contre les défauts de fabrication par un programme de garantie internationale. Pour plus d'informations sur la garantie, contactez le fabricant: Rossmax International Ltd. Attention: Consultez les documents d'accompagnement. Veuillez lire ce manuel attentivement avant l'emploi. Pour obtenir des informations spécifiques sur la pression artérielle, contactez votre médecin. Veuillez à conserver ce manuel.

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