Amplivox Ltd
Model 170 Automatic Audiometer
Operating Manual
(Appplies from serial number 22966 onwards)

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1 Introduction

Thank you for purchasing an Amplivox audiometer. The Amplivox Model 170 is an automatic screening audiometer that will give many years of reliable service if treated with care.

1.1 Intended applications

The Model 170 screening audiometer is designed for use by general practitioners, occupational health staff and child health professionals and is the ideal instrument for primary care groups, schools and industry. The audiometer may also be used for manual audiometry, but it is not intended for use to determine the full extent and scope of a patient’s hearing deficiency.

Test results may be printed using the specified printer option or transferred to a PC running the Amplivox Audibase or AudiView applications.

1.2 Unpacking

Open the shipping carton and carefully remove all the equipment. Check against the delivery note that all the accessories ordered have been included with your audiometer. If anything is missing, please contact Amplivox Customer Support (+44 1865 880846; sales@amplivox.ltd.uk). If you have purchased from a distributor you should contact them directly.

Please retain the shipping carton and packing materials as the audiometer will need calibrating on an annual basis and should be returned to Amplivox in its original shipping carton.

1.3 Warranty card (UK Customers Only)

Please complete the enclosed warranty registration card and return it to Amplivox. This will enable us to register your purchase, help with your enquiries and provide technical support.

1.4 Standard contents

Model 170 Audiometer
Carrying case
Mains adapter
Operating manual & Audiview
Audiometric headset
Patient response switch
Audiogram cards
Calibration certificate
1.5 Optional accessories

Additional audiogram cards
Audibase software        USB Cable
Printer(s)               Printer cable(s)
Audiocups (noise reducing earphone enclosures)

2 Important Safety Instructions

The Model 170 instrument must be used only by practitioners qualified to perform audiometric tests. It is intended for use as a screening tool.

2.1 Precautions

READ THIS OPERATING MANUAL BEFORE ATTEMPTING TO USE THE INSTRUMENT

To comply with the standards IEC 60601-1 for safety and IEC 60601-1-2 for EMC the audiometer is designed to be used only with the medically-approved mains adapter supplied, which is specified as part of the equipment. **Do not use any other type of mains adapter with this instrument.** Refer to Section 12 for the stock number of the adapter.

The audiometer is for indoor use only and should be used only as described in this manual.

The transducers supplied with the audiometer are specifically calibrated with it; if these transducers are changed calibration will be required.

Do not immerse the unit in any fluids. See Section 8 of this manual for the proper cleaning procedure for the instrument and its accessories.

Do not use the instrument in an oxygen-rich environment or in the presence of a flammable anaesthetic mixture or other flammable agents.

Do not drop or otherwise impact this instrument. If the instrument is dropped or damaged, return it to the manufacturer for repair and/or calibration. Do not use the instrument if any damage is suspected.
The instrument must be stored and used within the specified temperature, pressure and humidity ranges (see Sections 7 and 9).

Do not attempt to open, modify or service the instrument. Return the instrument to the manufacturer or distributor for all repair and servicing requirements. Opening the instrument will void the warranty.

2.2 Electromagnetic compatibility (EMC) considerations

Medical electrical equipment needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in Appendix 1. This provides guidance on the electromagnetic environment in which to operate the instrument.

Portable and mobile radio-frequency (RF) communications equipment can affect medical electrical equipment. The instrument should not be used adjacent to or stacked with other equipment; if this is necessary the instrument should be observed to verify normal operation.

2.3 Mains supply operation

The audiometer is designed for continuous operation and is powered by a mains adapter which is supplied, and specified as part of the equipment. If a replacement is required, please contact your Amplivox distributor.

All other connections must be made **before** connecting the output lead from the adapter into the POWER input socket on the back of the audiometer. Switch on the mains supply - the indicator on the adapter and the POWER indicator on the audiometer will both illuminate green, showing that the instrument is ready for use.

The output from mains adapter is fitted with electronic circuit protection. In case of overload the adapter will shut down and the indicator will be extinguished. When the fault is cleared the adapter will operate as normal.

The input to the mains adapter is protected with a non-replaceable fuse. If this fails the adapter will not operate.

The mains adapter is the mains disconnect device and therefore the audiometer should be positioned such that easy access to the mains adapter is possible.
2.4 Audiometer connections

All the relevant accessory terminals and connections are labelled to ensure correct identification and connection as follows:

<table>
<thead>
<tr>
<th>Socket Label</th>
<th>Socket Type</th>
<th>Colour Code</th>
<th>Connected Part</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>RIGHT</td>
<td>6.3mm jack</td>
<td>Red</td>
<td>Air conduction headset *</td>
<td></td>
</tr>
<tr>
<td>LEFT</td>
<td>6.3mm jack</td>
<td>Blue</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PRINTER</td>
<td>RJ12 socket (6-way)</td>
<td></td>
<td>Printer *</td>
<td>See 2.5</td>
</tr>
<tr>
<td>USB</td>
<td>USB Connector Type B</td>
<td></td>
<td>Computer (via USB port)</td>
<td>See 2.6</td>
</tr>
<tr>
<td>N/A</td>
<td>6 pin mini DIN</td>
<td></td>
<td>Reserved port; Amplivox</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>diagnostic use only</td>
<td></td>
</tr>
<tr>
<td>POWER</td>
<td>2.5mm power jack</td>
<td></td>
<td>Mains AC/DC Adapter *</td>
<td></td>
</tr>
<tr>
<td>RESPONSE</td>
<td>6.3mm jack</td>
<td>Black</td>
<td>Patient Response Switch *</td>
<td></td>
</tr>
</tbody>
</table>

The relevant part numbers are indicated in Section 12

**Note regarding the 6-pin mini DIN connector:**
This is a restricted socket for Amplivox use only. No user access is permitted.

For connected parts marked * only connect the accessories supplied with the instrument or supplied by Amplivox or an Amplivox distributor. These parts have been tested for use with the Model 170 Screening Audiometer for compliance with the standards IEC 60601-1 and IEC 60601-1-2. The use of accessories other than those specified may compromise compliance with these standards. For other sockets refer to Appendix 2.

2.5 Data transfer to a printer

Please refer to Appendix 2 for important information regarding the connection of non-medical electrical equipment to medical electrical equipment

The audiometer can be upgraded with an option to allow connection to one of two designated portable thermal printers for printing audiometric test results (see Section 3.8). You must use the designated cable for each printer, which is supplied with this option.
Upon receipt of the printer it must be initially charged for a **minimum of 15 hours** prior to use.

### 2.6 Data transfer to a computer

Please refer to Appendix 2 for important information regarding the connection of non-medical electrical equipment to medical electrical equipment.

The audiometer is supplied with software to allow connection to a computer for the transfer of test results (see Section 3.9). You must use the designated USB cable which is available from Amplivox (see Section 12).

### 3 Using the Audiometer

#### 3.1 Switching the audiometer on and off

Press and briefly hold the switch marked \( \mathcal{U} \) (located on the back panel). No warm-up time is required. The display will briefly show the model and the type of headphone currently in use.

The display will then be similar to that shown in Section 3.3.

To switch off, press and hold the MENU key followed by the YES (RIGHT) key and then release both.

#### 3.2 Testing the patient response switch

Press the patient response switch and the light labelled RESPONSE (above and to the right of the display) will illuminate green.

#### 3.3 Audiometer display

On start-up the display will show the following default setting:

<table>
<thead>
<tr>
<th>SIGNAL dBHL</th>
<th>FREQUENCY Hz</th>
</tr>
</thead>
<tbody>
<tr>
<td>30dB</td>
<td>1kHz</td>
</tr>
<tr>
<td>&lt; &gt;</td>
<td>&lt; &gt;</td>
</tr>
</tbody>
</table>
This indicates that when the PRESENT key is pressed, a tone will be presented at 30dBHL at a frequency of 1kHz (1000Hz) to the indicated ear. On start up the audiometer defaults to the left ear.

3.4 Audimeter controls

3.4.1 Multifunction Keys

Several keys on the audimeter have different functions depending on the actual mode of operation. These are MENU (OFF), PRINT (RESET), LEFT (NO), RIGHT (YES), AUTO (RESULTS) and FREQUENCY  (MENU SELECT). The use of these keys is described below.

3.4.2 MENU

Press and hold MENU to access the following options. Use the MENU SELECT keys to step through the available options and then the NO, YES or SIGNAL  keys to select an action or modify a setting. Release of the MENU key then initiates the action or saves the modified setting and returns to the default display.

<table>
<thead>
<tr>
<th>Menu Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Switch off?:</td>
<td>As described in Section 3.1</td>
</tr>
<tr>
<td>Clear test?:</td>
<td>Press YES and release MENU to clear the Threshold Retention Function results from the previous test</td>
</tr>
<tr>
<td>Save audiogram to (1):</td>
<td>Use the SIGNAL  keys to select the required storage location and press the YES key to save the audiogram; then release MENU</td>
</tr>
<tr>
<td>Load audiogram no (1):</td>
<td>Use the SIGNAL  keys to select the required storage location and press the YES key to load the audiogram; then release MENU</td>
</tr>
<tr>
<td>Contrast:</td>
<td>Adjust contrast using the SIGNAL  keys; then release MENU</td>
</tr>
<tr>
<td>Print audiogram?:</td>
<td>Press YES and release MENU; then press YES to confirm the print operation or NO to cancel</td>
</tr>
</tbody>
</table>
Use 250Hz in auto?: Press YES to include 250Hz in an automatic test or NO to exclude from the test

Use 1K5Hz in auto?: Press YES to include 1.5kHz in an automatic test or NO to exclude from the test

Use 8KHz in auto?: Press YES to include 8kHz in an automatic test or NO to exclude from the test

Use familiarization?: Press YES to employ a familiarisation routine at the start of an automatic test sequence on either ear (see Section 4.5.2)

Store on 2 of 3?: Press YES to activate automatic storage of the threshold level at which the patient makes 2 out of 3 responses in a manual test

Pulse in Manual?: Press the YES key to output a pulse tone in manual mode

2 of 3 in auto?: Press YES to activate automatic storage of the threshold level at which the patient makes 2 out of 3 responses in an automatic test (rather than the default 3 out of 5)

Default level: Use the SIGNAL \(\downarrow\uparrow\) keys to adjust the default tone presentation level in manual mode

Select printer: Use the MENU SELECT to select either the Able AP1300 or the Martel MCP8830 printer

3.4.3 Description of Function of Other Keys

PRINT Press this key to print the current threshold levels; then press YES to confirm the print operation or NO to cancel

RESET Press this key during an automatic test to cancel the test and return to the default display; any thresholds already found will be retained

+20dB This enables tone levels to be presented with up to 20dB higher output in manual test mode;
press the key and then use SIGNAL \( \uparrow \) to access the extra 20dB in 5dB steps; an indicator above the key is used to show that the function is active

**AUTO**
This initiates an automatic test on the indicated ear (see Section 4.5)

**RESULTS**
Use this key at the conclusion of an automatic test to view the results (see Section 4.5.4)

**TALK OVER**
Hold this key to interrupt the test and route the operator’s voice from the front panel microphone to the headset; the level is adjusted with the SIGNAL \( \downarrow \uparrow \) keys; if an automatic test is in progress the current test frequency will be retested from the default level when the TALKOVER key is released

**LEFT**
Press once to select the left ear (the indicator above the key illuminates green); if the left ear is already selected press again to store the displayed signal value as a threshold (see Section 3.5.1)

**RIGHT**
Press once to select the right ear (the indicator above the key illuminates green); if the right ear is already selected press again to store the displayed signal value as a threshold (see Section 3.5.1)

**SIGNAL**
Press the \( \downarrow \uparrow \) keys to decrease or increase the level of the tone presented in 5dB steps; to scroll through the range keep the key pressed

**FREQUENCY**
Press the \( \leftrightarrow \) key to select a lower frequency and the \( \Rightarrow \) key to select a higher frequency

**PRESENT**
Press to present the displayed test signal to the patient. The “PRESENT” indicator above the display will be illuminated green during tone presentation.
3.5 Threshold Retention Function

This function records the thresholds for both ears at each frequency tested. Thresholds may be recorded manually or automatically.

The operator can then review the results at the end of the test and record them on an audiogram card, print them with the optional printer (see Section 3.8), save them to the internal memory (see Section 3.6) or transfer the results to a computer (see Section 3.9).

3.5.1 Recording thresholds manually

Once a threshold has been determined press the “selected” ear key once again. The threshold will be recorded and displayed as shown in the illustration in Section 3.5.3. **Note: this function will not operate if the “Store on 2 of 3” option has been enabled (see Section 3.5.2).**

3.5.2 Recording thresholds automatically

If the “Store on 2 of 3” option has been enabled (see Section 3.4.2) then a threshold will be recorded automatically by the audiometer if the patient makes a response to two out of three manual tone presentations at the same level and frequency. Thresholds determined using the “Store on 2 of 3” option are displayed within square brackets.

3.5.3 Reviewing retained thresholds

To review the retained thresholds, select the required frequency using the FREQUENCY keys. The recorded values for the left and right ears are shown on the lower line of the display as illustrated below.

<table>
<thead>
<tr>
<th>SIGNAL dBHL</th>
<th>FREQUENCY Hz</th>
</tr>
</thead>
<tbody>
<tr>
<td>30dB</td>
<td>4kHz</td>
</tr>
<tr>
<td>20</td>
<td>10</td>
</tr>
</tbody>
</table>

This display shows thresholds at 4kHz
Left ear 20dBHL
Right ear 10dBHL

To clear the Threshold Retention memory, use the Clear Test menu option described in Section 3.4.2.
3.6 Saving audiograms in internal memory

The user may save up to 12 audiograms, referenced by number, in the internal memory of the audiometer. To save the current set of audiogram thresholds (these are the “retained” values described in Section 3.5) press and hold the MENU key, and then press MENU SELECT repeatedly until “Save Audiogram to 1” appears on screen. Use the SIGNAL keys to select a location numbered 1-12, and then press the YES key. Release the MENU key once confirmation appears on the display.

Note that the Save process will overwrite any records that exist in the selected memory location.

3.7 Loading audiograms from internal memory

Press and hold the MENU key, and then press MENU SELECT repeatedly until “Load Audiogram No 1” appears on screen. Use the SIGNAL keys to select a location numbered 1-12, and then press the YES key. Release the MENU key once confirmation appears on the display.

3.8 Printing audiograms

Two designated thermal printers (the Able AP1300 or the Martel MCP8830) are available as options for use with the Model 170 audiometer. The correct printer must be selected (use the MENU options described in Section 3.4.2 to make this selection).

- Connect the PRINTER socket on the audiometer (6-way RJ12) to the printer with the supplied cable (refer to Section 2.5 of this operating manual for printer set-up). Note that the printer cables for the Able printer (A108) and Martel printer (A107) are not compatible.
- Ensure the printer is fully charged, switched on, loaded with paper and ready to print.
- Load the desired audiogram as described in Section 3.7; to print the current audiogram ignore this instruction.
- Press the PRINT key and on the prompt “Is printer ready?” press the YES key. The audiogram will then print. To cancel the print operation press NO.

3.9 Data transfer to Audibase or AudiView

Test results stored within the audiometer may be transferred to the Amplivox Audibase database which is available as an option and must be
installed on to a computer (see Section 12 for the part number). Alternatively, Amplivox AudiView allows data to be transferred to a computer and subsequently viewed, annotated & printed. This software is supplied on a CD which includes this operating manual.

Refer to the installation & operating instructions provided with Audibase or AudiView for further details.

4 Suggested Sequence of Operation and Test Procedure

The following applies to air conduction measurements. For illustrative purposes 5dB steps are used. Refer also to ISO 8253 for guidance.

4.1 Audiometry preparation and ambient conditions

Refer to the appropriate audiometric standards and other relevant publications for guidance on audiometric testing.

Audiometric testing should always be performed in quiet conditions (e.g. a quiet room or an acoustic booth). The optional Audiocups can provide an additional level of isolation from ambient noise. For further explanation on permissible ambient noise levels, please refer to the standard ISO6189.

4.2 Test system arrangement

The schematic diagram below shows a typical example of the use of audiometric test equipment. The audiometer is located on the desk of a seated operator as shown.

The patient is seated in front of the desk facing away from the operator. The patient wears a headset (see Section 4.3) and responds to test stimuli by use of a hand-held switch which is also connected to the instrument.
4.3 Headset

The headset must be fitted by a qualified person to ensure a proper seal and a comfortable fit. The leads from the headset are connected to the instrument and the headset is then fitted to the patient.

4.4 Manual audiometry

4.4.1 Pre-test

(1) Switch the audiometer on
(2) Perform a listening check
(3) Decide whether to use the manual or automatic Threshold Retention Function and/or an audiogram card to record the thresholds
(4) If the automatic Threshold Retention Function is required ensure that the Store on 2 of 3 option is enabled (see Section 3.4.2) and that a patient response switch is in use
(5) Prepare the test environment & patient (see Sections 4.1 to 4.3)
(6) If the patient response switch is not being used give instructions to the patient to acknowledge any tone presented by raising or lowering the finger
(7) If the patient response switch is in use give instructions to the patient to acknowledge any tone presented as follows:

“As soon as you hear the tone, press the switch. When you no longer hear the tone, release the switch”.

(8) Fit the headset to the patient. Select the better hearing ear (according to the patient) by pressing either the LEFT or RIGHT key and start the familiarisation session.

4.4.2 Familiarisation

(1) Present the tone 30dB at 1kHz for between 1 and 2 seconds. If there is no response at 30dB, increase the attenuation level in 10dB steps until the patient responds
(2) When the patient responds, wait for 1 to 2 seconds and present the tone again at the same level; however, if the patient does respond at 30dB, reduce the signal level in 10dB steps, repeating the presentation until there is no response, then increase the signal level in 5dB steps until the patient responds; wait 1 to 2 seconds and present the tone again at the same level
(3) If the responses are consistent with the pattern of tone presentation proceed to Section 4.4.3 and start measuring the patient’s hearing thresholds; if not, repeat the familiarisation process

4.4.3 Test

(1) Use the Clear test option (see Section 3.4.2) to clear any thresholds
(2) Present the first test tone at 30dB at 1kHz
(3) If the patient responds, reduce the signal level in 10dB steps repeating the presentation until there is no response; then increase the signal level in 5dB steps until the patient responds
(4) If the patient does not respond, increase the signal level in 5dB steps until there is a response and then continue with step 4.
(5) Repeat the test by reducing the signal level in 10dB steps until the patient no longer responds. Then increase the signal level in 5dB steps until a response occurs and note this level.
(6) Repeat step 5 until the patient responds three out of a maximum of five times at the same signal level. This indicates the patient’s hearing threshold level for that frequency. Either mark the threshold on an audiogram card or press the appropriate ear key once to activate the Threshold Retention Function and save the threshold level on screen.
(7) Proceed to the next test frequency. It is common practice to test the frequencies in the following order: 1k, 2k, 3k, 4k, 6k, 8k and 500 Hz.
(8) Repeat steps 2 to 7 for the other ear.

4.4.4 Post-test

(1) Use the Threshold Retention Function to review the results (See Section 3.5)
(2) If required do one or more of the following:
   - Record the results on an audiogram card, or
   - Save the results to the internal memory (Section 3.6), or
   - Print the results (Section 3.8), or
   - Transfer the results to a computer (Section 3.9)

Refer to Section 3.4.2 to clear the thresholds at the end of a test and, if required, switch off the audiometer.
4.5 Automatic audiometry

4.5.1 Pre-test

(1) Switch the audiometer on
(2) Perform a listening check
(3) Use the following MENU options (see Section 3.4.2) to adjust the automatic test settings as required.

- Use 250Hz in auto?
- Use 1K5Hz in auto?
- Use 8KHz in auto?
- Use familiarisation?
- 2 of 3 in auto?

(4) Give the following instructions to the patient:

“As soon as you hear the tone, press and release the response switch.”

(5) Fit the headset to the patient.

4.5.2 Familiarisation

If the “Use familiarization” option has been selected (see Section 3.4.2) an automatic test will commence with a trial run at 1kHz starting from -10dB to allow the patient to become familiar with the increasing level and operating the response switch.

If the automatic familiarisation run is not used (or if the patient is having difficulty responding to the presented tones) the familiarisation process described in Section 4.4.2 may be used.

4.5.3 Test

(1) To test both ears ensure that the left ear is selected
(2) To test the right ear only ensure that the right ear is selected; to test the left ear only cancel the test by pressing the RESET key once the right ear testing has commenced
(3) To initiate a test press the AUTO key
(4) If selected the familiarisation option will run
(5) The automatic test proper will then commence
The test will proceed, starting at 1kHz followed by the higher frequencies before testing at lower frequencies. The frequencies 125Hz and 750Hz are always omitted from an automatic test, along with any other frequencies that have been specifically excluded (250Hz, 1.5kHz or 8kHz).

The test may be cancelled at any time by pressing the RESET key; any thresholds already established will be retained unless cleared or overwritten.

Automatic testing proceeds by increasing the tone level in 5dB steps until a response is made, then decreasing the level by 10dB and presenting another tone. If there is no response the level is increased in 5dB steps, and when a response is made the level is attenuated again by 10dB.

When 3 responses are made to 5 tone presentations at the same level (“3of5”) this is taken to be the threshold. The “2 of 3 in auto” option records a threshold if 2 responses are made to 3 tone presentations.

If an error occurs, for example the patient does not respond to the loudest tone presented or holds down the response switch continuously then the test will pause with a message displayed. Refer to Section 4.5.5.

The TALKOVER key may be used to interrupt the test and give further instructions to the patient (see Section 3.5.3).

An automatic test concludes with a re-test at 1kHz to ensure that consistent responses have been made. If the threshold levels for the two tests are within 10dB the threshold established for the re-test will be retained. Otherwise the test will pause with a message displayed. Refer to Section 4.5.5.

**4.5.4 Post-test**

When an automatic test has concluded the “Test finished” message is displayed. Pressing the RESULTS key will display the thresholds that were established (use the LEFT & RIGHT keys to select the required ear). Use the FREQUENCY keys to view all the frequencies. One or more of the following actions may then be taken:

- record the thresholds manually on an audiogram card
- print the results by pressing the PRINT key
- return to the default display by pressing the MENU key
If required the operator may then add or modify automatically-generated thresholds using manual audiometry (see Section 4.4)

The thresholds are retained by the audiometer and may be viewed, stored, printed or transferred to a computer as described in Section 4.4.4.

Refer to Section 3.4.2 to clear the thresholds at the end of a test and, if required, switch off the audiometer.

4.5.5 Error messages

Four error messages are possible while an automatic test is running. Depending on circumstances it may be necessary to provide further instruction to the patient and/or perform manual familiarisation (see Section 4.4.2).

No response!

This occurs when the patient has made no response and the tone level has reached the maximum value. An option to repeat the test at that frequency is presented. Press YES to repeat or NO to skip to the next frequency. If the frequency is skipped the message “Test finished incomplete” will be displayed at the conclusion of the test.

Response always!

This indicates that the patient has not released the response switch and the tone level has reached the minimum value. An option to repeat the test at that frequency is presented. Press YES to repeat or NO to skip to the next frequency. If the frequency is skipped the message “Test finished incomplete” will be displayed at the conclusion of the test.

1KHz match exceeded!

This occurs when the threshold level found at the 1kHz re-test differs by more than 10dB from that found for the 1st test (see Section 4.5.3). An option to repeat the re-test is presented. Press YES to repeat or NO to accept the threshold level found at the re-test.

Test finished incomplete

This occurs if the audiometer was unable to record a threshold at one or more frequencies (e.g. if no response was made and the retry option was
not chosen). The operator then has the option to use manual audiometry to obtain any missing thresholds.

5 Specification

5.1 Output data

Outputs: Left and Right earphone  
Frequency range: 125Hz-8kHz  
Frequency accuracy: <1%  
Distortion: <2%  
Output level range: -10dBHL minimum; see Section 5.2 for maximum  
Output level accuracy: Within 3dB  
Output level step size: 5dB  
Output transducer: DD45 earphones (supplied)  
Tone present: Single or pulsed  
Communication: Integral talk over facility  
USB interface: Transfer of test results to a computer

5.2 Maximum hearing levels provided at each frequency

<table>
<thead>
<tr>
<th>Frequency, Hz</th>
<th>Air conduction, dBHL (DD45)</th>
<th>Frequency, Hz</th>
<th>Air conduction, dBHL (DD45)</th>
</tr>
</thead>
<tbody>
<tr>
<td>125</td>
<td>80</td>
<td>2000</td>
<td>100</td>
</tr>
<tr>
<td>250</td>
<td>100</td>
<td>3000</td>
<td>100</td>
</tr>
<tr>
<td>500</td>
<td>100</td>
<td>4000</td>
<td>100</td>
</tr>
<tr>
<td>750</td>
<td>100</td>
<td>6000</td>
<td>100</td>
</tr>
<tr>
<td>1000</td>
<td>100</td>
<td>8000</td>
<td>100</td>
</tr>
<tr>
<td>1500</td>
<td>100</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

5.3 Physical Data

Display: 2 lines of 24 characters  
Mains power: 100-240Vac; 50/60Hz; 0.4A  
Dimensions: 270mm long x 175mm deep x 68mm high  
Weight: 0.75kg (approx)  
Safety: IEC 60601-1 (plus UL, CSA & EN deviations)  
EMC: IEC 60601-1-2  
CE mark: To the EU Medical Device Directive
5.4 Equipment classification

<table>
<thead>
<tr>
<th>Type of protection against electric shock</th>
<th>Powered via SELV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Degree of protection against electric shock</td>
<td>ClassII mains adapter</td>
</tr>
<tr>
<td>Degree of protection against ingress of water</td>
<td>Type B applied part</td>
</tr>
<tr>
<td>Mode of operation</td>
<td>Not protected</td>
</tr>
<tr>
<td>Equipment mobility</td>
<td>Continuous operation</td>
</tr>
</tbody>
</table>

The Model 170 Audiometer is classified as a Class Ila device under Annex IX of the EU Medical Devices Directive. It is intended for use as a screening audiometer instrument.

6 Symbols

The following symbols appear on the audiometer or mains adapter:

Definition: Refer to instruction manual (mandatory).

Definition: Type B applied part – an applied part providing protection against electric shock, particularly regarding allowable patient leakage current and patient auxiliary current.

The applied parts are the left & right earphones, patient response switch and the associated cables.

Definition: The output from the mains AC adapter is Direct Current.

Definition: Class II equipment – equipment in which protection against electric shock does not rely on basic insulation only, but in which additional safety precautions such as double insulation or reinforced insulation are provided, there being no provision for protective earthing or reliance upon installation conditions.
7 Technical Information

Audiometer
Audiometer type: Type 4 (IEC 60645-1:2001)
Type 4 (ANSI S3.6:2004)

Transducers
Types and reference levels: DD45: ISO 389-1, Table 2
Static headband force: Headphones: 4.5N
Sound attenuation characteristics: ISO8253-1, Table 3

Earphone Sound Attenuation Characteristics

<table>
<thead>
<tr>
<th>Frequency, Hz</th>
<th>125</th>
<th>250</th>
<th>500</th>
<th>1000</th>
<th>2000</th>
<th>4000</th>
<th>8000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Attenuation, dB</td>
<td>3</td>
<td>5</td>
<td>7</td>
<td>15</td>
<td>26</td>
<td>32</td>
<td>24</td>
</tr>
</tbody>
</table>

Environmental
Operating temperature: +15°C to +35°C
Operating humidity: 30% to 90% (non-condensing)
Atmospheric pressure: 700 hPa to 1060 hPa

Input / Output
Power input: 2.5mm barrel-type socket.
Patient response input: 6.3mm Jack socket
Left & Right outputs: 6.3mm Jack socket
USB: Type B socket
Printer: RJ12 socket (6-way)
Maximum voltage at any output: 12V peak

8 Routine Maintenance

8.1 Audiometer maintenance

The Model 170 audiometer is a precision instrument. Handle it carefully in order to ensure its continued accuracy and service. When cleaning the instrument, first disconnect it from the mains supply. Use a soft cloth and mild detergent to clean the instrument panel when required. Refer to ISO 8253-1 for additional guidance.

8.2 Transducer maintenance

Before use check the transducer cables and connectors for signs of wear and/or damage. If you find any, please replace the item immediately by
contacting Amplivox or your Amplivox distributor, requesting the relevant part number (see Section 12).

Handle the audiometric headset and other accessories with care. For parts that are in direct contact with the patient it is recommended that replacement parts are used or the parts are subjected to a standard disinfecting procedure between patients.

This includes physically cleaning and use of a recognised disinfectant. The specific manufacturer’s instructions should be followed for use of this disinfecting agent to provide an appropriate level of cleanliness.

Clean the ear cushions (including those on the Audiocups, if used) with a recognised disinfectant, e.g. a “Mediswab”.

![Warning]

During the cleaning process do not allow moisture to enter the earphone.

8.3 Mains adapter maintenance

Before use check the mains AC adapter for signs of wear and/or damage. If you find any replace the adapter immediately by contacting Amplivox or your Amplivox distributor. Refer to Section 12 for approved part numbers.

![Warning]

DO NOT USE ANY OTHER TYPE OF MAINS ADAPTER WITH THIS INSTRUMENT. See Section 2.3.

9 Instrument Storage and Transportation

This instrument can be stored or transported within the following environmental parameters:

Temperature: -20°C to +70°C
Humidity: 10% to 90% (non-condensing)
Atmospheric Pressure: 500 hPa to 1060 hPa
10 Calibration and Repair of the Instrument

Amplivox recommends that this audiometer should be calibrated on an annual basis. Please contact Amplivox or the designated distributor for details of calibration services. Refer to ISO 8253-1 for additional guidance.

The instrument should be returned to the manufacturer for service & repair. There are no user-serviceable parts within it.

When packing the instrument for shipping, please use the original shipping carton and packing materials. Please also ensure that the headset leads are not wrapped around the headband of the headset.

11 Guarantee

All Amplivox instruments are guaranteed against faulty materials and manufacture. The instrument will be repaired free of charge for a period of two years from the date of despatch if returned, carriage paid, to the Amplivox service department. Return carriage is free of charge for customers in the UK and chargeable for overseas customers.

Important Note:
The following exceptions apply:

Earphones may go out of calibration due to rough handling or impact (dropping). The life of the leads is also dependent upon conditions of use. These parts are only guaranteed against faulty materials or manufacture.
12 Ordering Consumables and Accessories

To order consumables, additional accessories and to replace detachable parts that have been damaged, please contact Amplivox for current prices and delivery charges. The items available are listed below:

**Stock No.**  **Description**

A022  Audiocups (noise reducing earphone enclosures)
AC1042  Audiocup ear cushion
AC1047  Audiocup headband
AC1048  Audiocup headband cover
A023  Headband (standard headphone)
A026  Earphone cushion
A032  Earphones DD45 *
A030  Headset lead

B128  Carrying case
A091-7  Approved mains adapter
A085  Patient response switch
A051  Audiogram cards (pack of 50)

A091  Printer Martel MCP8830
A107  Printer cable for audiometer to Martel MCP8830
C01  Thermal Printer paper for Martel MCP8830
PT01  Printer Able AP1300
A108  Printer cable for audiometer to Able AP1300
C0103  Thermal Printer paper for Able AP1300

F07  USB Cable, 1.8m
AUD06  Amplivox Audibase 5.5 (including USB cable)

**Accessories marked * require calibration with the specific audiometer to be used. Do not attempt to use these accessories until the audiometer has been calibrated to match their characteristics.**

Shipping documentation will reference the stock number quoted above, and images of the parts alongside the relevant stock number are available on the Amplivox website (www.amplivox.ltd.uk). The required fitting instructions are supplied with each part.
13 Disposal Information

Amplivox Limited is fully compliant with the WEEE (Waste Electrical and Electronic Equipment) Regulations. Our PRN (Producer Registration Number) is WEE/GA0116XU and we are registered with the approved WEEE Compliance Scheme, B2B Compliance, approval number WEE/MP3338PT/SCH.

The main purpose of the WEEE Regulations is to encourage the segregation of waste electrical items from the general waste stream and into reuse, recovery and recycling routes.

For any waste electrical units purchased from Amplivox that either:

- bear the crossed out wheeled bin symbol with black bar underneath
- or, have been replaced with new Amplivox products on a like-for-like basis

please contact our WEEE Compliance Scheme using the details below. B2B Compliance will be able to provide further information on how to recycle your waste electrical units and answer any queries you may have.

**B2B Compliance**
Tel: +44 (0) 1691 676 124 (Option 2)
Email: operations@b2bcompliance.org.uk
Appendix 1 - EMC Guidance & Manufacturer’s Declaration

<table>
<thead>
<tr>
<th>Emissions test</th>
<th>Compliance</th>
<th>Electromagnetic environment – guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td>Group 1</td>
<td>The Model 170 Audiometer uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RF emissions</td>
<td>Class A</td>
<td>The Model 170 Audiometer is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td>Class A</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations/flicker emissions</td>
<td>Complies</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Guidance and manufacturer’s declaration – electromagnetic immunity (1)

The Model 170 Audiometer is intended for use in the electromagnetic environment specified below. The customer or user of the Model 170 Audiometer should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment – guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic Discharge (ESD)</td>
<td>±6 kV contact 8 kV air</td>
<td>±6 kV contact 8 kV air</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%</td>
</tr>
<tr>
<td>IEC 61000-4-2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Electrical fast transient/burst</td>
<td>±2 kV for power supply lines</td>
<td>±2 kV for power supply lines</td>
<td>Mains power quality should be that of a typical commercial or hospital environment</td>
</tr>
<tr>
<td>IEC 61000-4-4</td>
<td>±1 kV for input/output lines</td>
<td>±1 kV for input/output lines</td>
<td></td>
</tr>
<tr>
<td>Surge</td>
<td>±1 kV differential mode</td>
<td>±1 kV differential mode</td>
<td>Mains power quality should be that of a typical commercial or hospital environment</td>
</tr>
<tr>
<td>IEC 61000-4-5</td>
<td>±2 kV common mode</td>
<td>±2 kV common mode</td>
<td></td>
</tr>
<tr>
<td>Immunity test</td>
<td>IEC 60601 test level</td>
<td>Compliance level</td>
<td>Electromagnetic environment – guidance</td>
</tr>
<tr>
<td>------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines</td>
<td>&lt;5% U_T (&gt;95% dip in U_T) for 0.5 cycle</td>
<td>&lt;5% U_T (&gt;95% dip in U_T) for 0.5 cycle</td>
<td>Mains power quality should be that of a typical commercial or hospital environment. If the user of the Model 170 Audiometer requires continued operation during power mains interruptions, it is recommended that the Model 170 Audiometer be powered from an uninterruptible power supply or a battery</td>
</tr>
<tr>
<td>IEC 61000-4-11</td>
<td>40% U_T (60% dip in U_T) for 5 cycles</td>
<td>40% U_T (60% dip in U_T) for 5 cycles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>70% U_T (30% dip in U_T) for 25 cycles</td>
<td>70% U_T (30% dip in U_T) for 25 cycles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt;5% U_T (&gt;95% dip in U_T) for 5 sec</td>
<td>&lt;5% U_T (&gt;95% dip in U_T) for 5 sec</td>
<td></td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-8</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NOTE U_T is the a.c. mains voltage prior to the application of the test level
**Guidance and manufacturer’s declaration – electromagnetic immunity (2)**

The Model 170 Audiometer is intended for use in the electromagnetic environment specified below. The customer or user of the Model 170 Audiometer should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment – guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>IEC 61000-4-6</td>
<td>3 Vrms</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Model 170 Audiometer, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
<tr>
<td></td>
<td>150kHz to 80MHz</td>
<td>3 Vrms</td>
<td>d = 1.2√P 80MHz to 800MHz</td>
</tr>
<tr>
<td>Radiated RF</td>
<td>IEC 61000-4-3</td>
<td>3 V/m</td>
<td>Recommended separation distance</td>
</tr>
<tr>
<td></td>
<td>80MHz to 2.5GHz</td>
<td>3 V/m</td>
<td>d = 1.2√P 800MHz to 2.5GHz</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>d = 2.3√P 800MHz to 2.5GHz</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>where P is the maximum output power rating of the transmitter in Watts (W) according to the transmitter manufacturer and d is the recommended separation distance in metres (m).</td>
</tr>
</tbody>
</table>
### Guidance and manufacturer's declaration – electromagnetic immunity (2)

Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, \(^a\) should be less than the compliance level in each frequency range. \(^b\)

Interference may occur in the vicinity of equipment marked with the following symbol:

![Symbol](image)

**NOTE 1** At 80MHz and 800MHz, the higher frequency range applies.

**NOTE 2** These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

| a | Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephone, land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the Model 170 Audiometer is used exceeds the applicable RF compliance level above, the Model 170 Audiometer should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the Model 170 Audiometer. |
| b | Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m. |
Recommended separation distances between portable and mobile RF communications equipment and the Model 170 Audiometer

The Model 170 Audiometer is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Model 170 Audiometer can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Model 170 Audiometer as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter</th>
<th>Separation distance according to frequency of transmitter</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150 kHz to 80 MHz</td>
</tr>
<tr>
<td>W</td>
<td>m</td>
</tr>
<tr>
<td>------------------------------------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.38</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.8</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance $d$ in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where $P$ is the maximum output power rating of the transmitter in Watts (W) according to the transmitter manufacturer.

NOTE 1 At 80MHz and 800MHz, the separation distance for the higher frequency range applies.

NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.
Appendix 2 - Use with Non-medical Electrical Equipment

Any person who connects external equipment to signal input, signal output or other connectors has created a medical electrical system and is therefore responsible for the system complying with the requirements of clause 16 of IEC 60601-1:2005 (General requirements for basic safety and essential performance).

If connections are made to standard equipment such as printers and computers, special precautions must be taken in order to maintain medical safety. The following notes are provided for guidance in making such connections to ensure that the general requirements of clause 16 of IEC 60601-1:2005 are met.

The following signal inputs and outputs on the Model 170 audiometer are electrically isolated to the requirements of IEC 60601-1 in order to reduce any potential hazard associated with the use of mains-powered equipment connected to these inputs and outputs:

<table>
<thead>
<tr>
<th>Socket Label</th>
<th>Socket Type</th>
<th>Typical Connection</th>
</tr>
</thead>
<tbody>
<tr>
<td>PRINTER</td>
<td>RJ12 socket (6-way)</td>
<td>Printer</td>
</tr>
<tr>
<td>USB</td>
<td>USB Connector Type B</td>
<td>Computer</td>
</tr>
</tbody>
</table>

External equipment intended for connection to signal input, signal output or other connectors, shall comply with the relevant IEC or international standards (e.g. IEC 60950, CISPR 22 & CISPR 24 for IT equipment, and the IEC 60601 series for medical electrical equipment).

Equipment not complying with IEC 60601 shall be kept outside the patient environment, as defined in IEC 60601-1:2005 (at least 1.5m from the patient).

The operator must not touch the connected equipment and the patient at the same time as this would result in an unacceptable hazard.

Refer to Diagrams 1 to 3 below for typical configurations of connected peripheral equipment. Refer to Amplivox Limited at the address given on the front of this user manual if advice is required regarding the use of peripheral equipment.
Diagram 1: Model 170 used with the medically-approved mains adapter

Mains Outlet  Medical Mains Adapter

Model 170 Audiometer
Diagram 2: Model 170 used with the medically-approved mains adapter and printer

Mains Outlet

Medical Mains Adapter

Model 170 Audiometer

Mains Outlet

Printer via PRINTER socket

Printer Power Supply
Diagram 3: Model 170 used with the medically-approved mains adapter and PC

- Mains Outlet
- Medical Mains Adapter
- Model 170 Audiometer
- PC Power Supply
- PC via USB socket