Warnings & Cautions

Reichert Technologies is not responsible for the safety and reliability of this instrument when:

- Assembly, disassembly, repair, or modification is made by unauthorized dealers or persons.
- Instrument is not used in accordance with this User’s Guide.

**WARNING:** AN INSTRUCTION THAT DRAWS ATTENTION TO RISK OF INJURY OR DEATH.

**WARNING:** UNITED STATES FEDERAL LAW AND EUROPEAN REGULATIONS REQUIRE THAT THIS DEVICE BE PURCHASED ONLY BY A PHYSICIAN OR A PERSON ACTING ON BEHALF OF A PHYSICIAN.

**WARNING:** THIS INSTRUMENT SHOULD BE USED IN STRICT ACCORDANCE WITH THE INSTRUCTIONS OUTLINED IN THIS USER’S GUIDE. THE SAFETY OF THE OPERATOR AND THE PERFORMANCE OF THE INSTRUMENT CANNOT BE GUARANTEED IF USED IN A MANNER NOT SPECIFIED BY REICHERT TECHNOLOGIES.

**WARNING:** DO NOT REPAIR OR SERVICE THIS INSTRUMENT WITHOUT AUTHORIZATION FROM THE MANUFACTURER. ANY REPAIR OR SERVICE TO THIS INSTRUMENT MUST BE PERFORMED BY EXPERIENCED PERSONNEL OR DEALERS WHO ARE TRAINED BY REICHERT SO THAT CORRECT OPERATION OF THIS INSTRUMENT IS MAINTAINED OR SERIOUS INJURY TO THE OPERATOR OR PATIENT MAY OCCUR.

**WARNING:** MODIFICATIONS TO THIS INSTRUMENT IS NOT ALLOWED. ANY MODIFICATION TO THIS UNIT MUST BE AUTHORIZED BY REICHERT SO THAT CORRECT OPERATION IS MAINTAINED OR SERIOUS INJURY TO THE OPERATOR OR PATIENT MAY OCCUR.

**WARNING:** IF THIS INSTRUMENT IS MODIFIED, APPROPRIATE INSPECTION AND TESTING MUST BE CONDUCTED TO ENSURE CONTINUED SAFE USE OF THIS INSTRUMENT.

**WARNING:** TO AVOID RISK OF ELECTRIC SHOCK, THIS EQUIPMENT MUST ONLY BE CONNECTED TO A SUPPLY MAINS WITH PROTECTIVE EARTH OR DAMAGE TO THE INSTRUMENT AND/OR INJURY TO THE OPERATOR OR PATIENT MAY OCCUR.

**WARNING:** ENSURE THAT THE VOLTAGE APPLIED TO THE UNIT IS THE SAME AS THE VOLTAGE THAT IS INDICATED ON THE DATA PLATE OR DAMAGE TO THE INSTRUMENT AND/OR INJURY TO THE OPERATOR OR PATIENT MAY OCCUR.

**WARNING:** THIS INSTRUMENT MUST BE PLUGGED INTO AN OUTLET WITH AN EARTH GROUND. DO NOT REMOVE OR DEFEAT THE EARTH GROUND CONNECTION ON POWER INPUT CONNECTOR OR THE UNIT’S POWER CORD OF THIS INSTRUMENT OR DAMAGE TO IT AND/OR INJURY TO THE OPERATOR OR PATIENT MAY OCCUR.

**WARNING:** THE EQUIPMENT OR SYSTEM SHOULD NOT BE USED ADJACENT TO OR STACKED WITH OTHER EQUIPMENT AND THAT IF ADJACENT OR STACKED USE IS NECESSARY, THE EQUIPMENT OR SYSTEM SHOULD BE OBSERVED TO VERIFY NORMAL OPERATION IN THE CONFIGURATION IN WHICH IT WILL BE USED.

**WARNING:** THIS INSTRUMENT IS NOT SUITABLE FOR USE IN THE PRESENCE OF FLAMMABLE ANESTHETIC MIXTURES, SUCH AS OXYGEN OR NITROUS OXIDE.

**WARNING:** DO NOT USE ANY OTHER PROBE WITH THIS INSTRUMENT THAN THOSE SUPPLIED BY REICHERT EXPRESSLY FOR USE WITH THIS INSTRUMENT. OTHER PROBES MAY CAUSE DAMAGE OR INJURY TO THE EYE.

**WARNING:** DO NOT USE THIS INSTRUMENT IF THE MEASUREMENT TIP IS CRACKED, CHIPPED OR SHOWS ANY IRREGULARITY OF THE SURFACE, TO PREVENT PATIENT INJURY AND OR INACCURATE READINGS.

**WARNING:** IN ORDER TO PREVENT PATIENT-TO-PATIENT TRANSFER OF INFECTION, AFTER EACH USE DISINFECT THE MEASUREMENT TIP FOLLOWING ACCEPTED LOCAL CLINICAL PROCEDURES REGARDING THE USE OF DISINFECTANTS. ANY CLINICALLY APPROVED CHEMICAL DISINFECTANT CAN BE USED.
WARNING: THE USE OF ACCESSORIES OR CABLES OTHER THAN THOSE SPECIFIED, WITH THE EXCEPTION OF THOSE SOLD BY THE MANUFACTURER AS REPLACEMENT PARTS FOR INTERNAL COMPONENTS, MAY RESULT IN INCREASED EMISSIONS OR DECREASED IMMUNITY OF THE EQUIPMENT OR SYSTEM.

CAUTION: AN INSTRUCTION THAT DRAWS ATTENTION TO THE RISK OF DAMAGE TO THE PRODUCT.

⚠️ CAUTION: THE INTERNAL CIRCUITRY OF THE INSTRUMENT CONTAINS ELECTROSTATIC DISCHARGE SENSITIVE DEVICES (ESDS) THAT MAY BE SENSITIVE TO STATIC CHARGES PRODUCED BY THE HUMAN BODY. DO NOT REMOVE THE COVERS WITHOUT TAKING PROPER PRECAUTIONS.

CAUTION: DO NOT USE SOLVENTS OR STRONG CLEANING SOLUTIONS ON ANY PART OF THIS INSTRUMENT AS DAMAGE TO THE UNIT MAY OCCUR. SEE MAINTENANCE SECTION FOR DETAILED CLEANING INSTRUCTION.

CAUTION: USE OF AMMONIA BASED CLEANERS ON THE LIQUID CRYSTAL DISPLAY (LCD) MAY CAUSE DAMAGE TO DISPLAY. SEE MAINTENANCE SECTION FOR DETAILED CLEANING INSTRUCTION.

CAUTION: DO NOT AUTOCLAVE OR DISINFECT USING HIGH TEMPERATURES EXCEEDING THE RECOMMENDED TEMPERATURES INDICATED IN THE SPECIFICATIONS SECTION OF THIS MANUAL OR DAMAGE TO THE UNIT MAY OCCUR.

CAUTION: MEDICAL ELECTRONIC EQUIPMENT NEEDS SPECIAL PRECAUTIONS REGARDING EMC AND NEEDS TO BE INSTALLED AND PUT INTO SERVICE ACCORDING TO THE EMC INFORMATION PROVIDED IN THE ACCOMPANYING DOCUMENTS.

CAUTION: PORTABLE AND MOBILE RF COMMUNICATIONS EQUIPMENT CAN AFFECT MEDICAL ELECTRICAL EQUIPMENT.

CAUTION: THIS INSTRUMENT IS NOT TO BE USED NEAR HIGH-FREQUENCY EMITTING SURGICAL EQUIPMENT.

CAUTION: THIS INSTRUMENT IS NOT INTENDED TO BE CONNECTED TO EQUIPMENT OUTSIDE THE CONTROL OF REICHERT INC. OR MUST BE TESTED TO AN APPLICABLE IEC OR ISO STANDARDS.

CAUTION: DO NOT INSTALL ANY ADDITIONAL SOFTWARE OTHER THAN WHAT WAS SUPPLIED WITH THIS INSTRUMENT. INSTALLATION OF ADDITIONAL SOFTWARE MAY CAUSE UNEXPECTED OPERATION RESULTING IN MALFUNCTION OF THIS INSTRUMENT.
Symbol Information

The following symbols appear on the instrument:

- Caution
- Type B Applied Part
- Alternating Current Power
- Protective Earth Connection
- ON / OFF
- Manufacturer
- Date of Manufacture
- REF Catalog Number
- S/N Serial Number
- Waste of Electrical and Electronic Equipment
- Compliance to Medical Device Directive 93/42/EEC
- Authorized to mark given by Intertek ETL Semko for conformance with electrical standards
- Accompanying Documents must be consulted
- Authorized Representative in European Community
- Fragile Contents in Shipping Container - handle with care
- Keep Dry - Package shall be kept away from rain
- This Way Up - Indicates correct upright position of package
- Important Instruction - remove Shipping Bracket
Congratulations on your purchase of the Reichert® Model 30™ Pneumatonometer.

The Model 30 Pneumatonometer is a highly accurate instrument used to measure intraocular pressure (IOP) non-invasively through applanation tonometry and represents a significant advance in tonometry and tonography technology over its predecessors. The pneumatonometer system consists of the base unit, a probe, and the necessary accessories to accurately measure tonometry/tonography in one easy-to-use unit. This manual describes its operation and details. Most individuals will find this instrument easy to use following the software and audible prompts.

This User’s Guide is designed as a training and reference manual for operation, maintenance, and troubleshooting. We recommend that you read it carefully prior to use and follow the instructions in the guide to ensure optimum performance of your new instrument. Properly trained eyecare professionals such as ophthalmologists, optometrists, opticians and eye care technicians should operate this instrument.

Please retain this guide for future reference and to share with other users. Additional copies can be obtained from your authorized Reichert Technologies dealer or contact our Customer Service department directly at:

Tel: 716-686-4500  
Fax: 716-686-4555  
Email: reichert.information@ametek.com

Please indicate the following when contacting Reichert to ensure that you receive the correct information:

- Model Number
- Serial Number
- Contact Phone Number or Email Address

**Indications for use**
The Model 30 Classic Pneumatonometer is intended for the measurement of intraocular pressure. The Model 30 Classic Pneumatonometer is indicated for use as a screening / monitoring tool for glaucoma or when increased intraocular pressure is suspected.

**Contraindications**
None.
Instrument Setup

Great care has been taken to deliver this instrument to you safely. The container and packaging was specially designed to transport this unit. Please retain the packaging if future transportation is required. Please remove the packaging material from the outer box and then remove the pneumatonometer and its accessories from the box. Refer to Figures 1-1 through 1-3.

Unpacking Instructions

Please remove the packaging material from the instrument in the following manner. Refer to Figures 1-1 through 1-3.

The instrument is packaged in a shipping container to protect the instrument from damage during shipment. Please read the User’s Guide before operating the unit. The pneumatonometer includes the following accessories:

1. User’s Guide (not shown)  
   (P/N 16030-101)
2. Spare Tip and Membrane Assembly  
   (P/N 230676)
3. Pneumatonometer Probe  
   (P/N 232349)
4. Power Cord  
   (P/N WCBL10018 for 16030 & 16032  
    or P/N WCBL10027 for 16031)
5. Footswitch  
   (P/N 232345 - only available with P/N 16033 kit)
6. Thermal Paper  
   (P/N 232348)
7. Calibration Verifier  
   (P/N 232373)
8. Yellow Shipping Bracket (not shown)  
   (P/N 16030-009)  
   (attached to the unit when shipped).
Parts Identification
The Model 30 Pneumatonometer is housed in a single compact metal case that fits easily on most counters or an appropriate stand.

The front panel of the Model 30 Pneumatonometer includes: (Refer to the Figure 1-4.)

1. ON/OFF switch
2. Built-in chart
3. LCD readout

The rear panel of the Model 30 Pneumatonometer (Figure 1-5) includes:

1. Access door for changing filters and removing the yellow shipping bracket.
2. Receptacle for the optional footswitch, which allows the operator to select system functions while using his/her hands to restrain the patient’s eyelid and position the probe.
3. Three-prong plug for the AC power cord that connects the Model 30 Pneumatonometer to an electrical outlet with an integral fuse holder.
4. USB connection for factory data communication.

Note: The USB connection is only for use by the manufacturer, for service repairs. This port cannot be used to transfer data or connect to an external printer.
Parts Identification

Models and Options
The specifications for the different models of the pneumatonometer are found in the General Specifications section of this manual.

The pneumatonometer is a tonometer that provides the following tests: manual tonometry, pulse tonometry or tonography.

In the manual mode, the pneumatonometer provides a single pressure value and ends the test when the probe is removed from the surface of the patient’s eye. The pulse mode provides pressure output with pulse fluctuations and ends the test after it has read ten ocular pulses or the probe is removed from the eye surface. The tonography test records changes in IOP when a weighted probe is held on the cornea for a specified time. The tonography test enables the operator to conduct a two-minute or four-minute test for each eye.

The Model 30 Pneumatonometer handpiece and probe assembly (Figure 1-6) includes:
1. Tip and membrane assembly (Applied Part)
2. Auto sensor assembly and probe handle
3. Double lumen tubing
4. Quick-flow connector

Accessories (Optional)

<table>
<thead>
<tr>
<th>Part Number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>16033</td>
<td>Tonography Kit (includes Foot Pedal &amp; 10 Gram Weight)</td>
</tr>
<tr>
<td>230676</td>
<td>Tip and Membrane Assembly, 1 pk</td>
</tr>
<tr>
<td>230677</td>
<td>Tip and Membrane Assembly, 3 pk</td>
</tr>
<tr>
<td>230678</td>
<td>Tip and Membrane Assembly, 10 pk</td>
</tr>
<tr>
<td>232346</td>
<td>Filter Kit</td>
</tr>
<tr>
<td>232348</td>
<td>Printer Paper</td>
</tr>
<tr>
<td>232373</td>
<td>Calibration Verifier</td>
</tr>
</tbody>
</table>

(To order any of these accessories, contact your local authorized Reichert dealer.)
**Icon Definition**

The Pneumatonometer incorporates a user-friendly icon/menu-based operating system that will increase the speed of measurements, training and use. Below are the Icons that are used during the operation of this instrument.

<table>
<thead>
<tr>
<th>Icon</th>
<th>Icon Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SETUP</td>
<td>Access the setup menu for changing default parameters.</td>
</tr>
<tr>
<td>MEASURE</td>
<td>Initiates the measurement process.</td>
</tr>
<tr>
<td>MANUAL MODE</td>
<td>Initiates the manual measurement mode process.</td>
</tr>
<tr>
<td>PULSE MODE</td>
<td>Initiates the pulse measurement mode process.</td>
</tr>
<tr>
<td>TONOGRAPHY</td>
<td>Initiates the tonography mode.</td>
</tr>
<tr>
<td>LEFT EYE</td>
<td>Left eye measurement.</td>
</tr>
<tr>
<td>RIGHT EYE</td>
<td>Right eye measurement.</td>
</tr>
<tr>
<td>PAPERFEED</td>
<td>Advances the printer paper.</td>
</tr>
<tr>
<td>2 MINUTE</td>
<td>Initiates the two minute measurement mode.</td>
</tr>
<tr>
<td>4 MINUTE</td>
<td>Initiates the four minute measurement mode.</td>
</tr>
<tr>
<td>ENTER DATA</td>
<td>Allows data to be entered manually.</td>
</tr>
<tr>
<td>CALCULATE</td>
<td>Provides calculation of data.</td>
</tr>
</tbody>
</table>
## Instrument Setup  (continued)

### Icon Definition (continued)

<table>
<thead>
<tr>
<th>Icon</th>
<th>Icon Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>RETURN</td>
<td>Returns to preceding screen.</td>
</tr>
<tr>
<td>RIGHT ARROW</td>
<td>Used in the setup menus to move right horizontally.</td>
</tr>
<tr>
<td>LEFT ARROW</td>
<td>Used in the setup menus to move left horizontally.</td>
</tr>
<tr>
<td>UP ARROW</td>
<td>Used in the setup menus to move up vertically.</td>
</tr>
<tr>
<td>DOWN ARROW</td>
<td>Used in the setup menus to move down vertically.</td>
</tr>
<tr>
<td>PLUS</td>
<td>Increases a numerical value displayed on the screen.</td>
</tr>
<tr>
<td>MINUS</td>
<td>Decreases a numerical value displayed on the screen.</td>
</tr>
<tr>
<td>SELECT</td>
<td>Used in the setup menus to activate the new parameter or setting.</td>
</tr>
<tr>
<td>HOME</td>
<td>Changes the display to the main menu screen.</td>
</tr>
<tr>
<td>PRINT</td>
<td>Provides a printout of the current data.</td>
</tr>
</tbody>
</table>
Default Settings

This instrument is sent from the factory with measurement, chart, and other general parameters assigned to default settings. These settings can be changed to the preferences of the operator/clinician. A summary of these settings is given below. The optional settings follow on subsequent pages.

General Setup:
- Chart: On
- Footswitch: Off
- Volume: Middle
- Brightness: Middle

Printout Setup:
- Date: (current date)
- Date Fmt: MDY
- Time: (current time)
- Time Fmt: AM/PM

Service:
- Service Screen with:
  - Software Version
  - Calibration Date
  - Customer Service Telephone Number

General Setup

Parameters:  Settings:
Chart: Options are either On or Off.
Footswitch: Options are either On or Off.
Volume: This option is a left or right arrow to either decrease or increase the volume.
Brightness: This option is a left or right arrow to either decrease or increase the brightness.
Instrument Setup  (continued)

Printout Setup

Parameters:  Settings:
Date: When the unit is received, the date and time should be changed to the local time zone.
DateFmt: The options are: MDY, DMY, YMD (D=day, M=month, Y=year)
Time: The correct time should be changed to agree with the local time zone.
TimeFmt: The options are: AM/PM, 24HR

Service Screen

This sample screen provides the details of the Software Version, Calibration Date, and the customer service telephone number. If assistance is needed with this unit, consult the Troubleshooting section of this manual. If further assistance is needed, please contact the customer service department at Reichert, Inc.
Instrument Setup (continued)

Installation

Consider the following factors as you find a location for the Model 30 Pneumatonometer:

• When taking tonometry measurements with the pneumatonometer, the patient can be either seated or supine. When conducting tonography tests, the patient must be supine.
• The console should be close enough to an electrical outlet so that the operator can connect the system with the power cord.
• The operator should be in a position to look directly at the contact between the probe and the surface of the eye.
• The operator should be able to see the instrument’s LCD readout and chart recorder easily during a test.
• This medical device complies with IEC/EN 60601-1-2 Safety Standard for Electromagnetic Compatibility, Requirements and Test. However, if the equipment is operated in the presence of high levels of EMI or highly sensitive equipment, interference may be encountered and the user should take whatever steps are necessary to eliminate or reduce the interference.

Pneumatonometer Installation

**WARNING:** CARE MUST BE TAKEN TO ARRANGE THE CABLES FOR THE ACCESSORIES SUCH THAT THEY DO NOT PRESENT A TRIPPING HAZARD TO THE EXAMINER OR A DANGER TO THE PATIENT.

**WARNING:** POSITION THIS INSTRUMENT SO THAT IT IS NOT DIFFICULT TO OPERATE THE DISCONNECTION DEVICE (PLUG).

1. Set the ON/OFF switch on the front panel to the OFF position (the O is pushed in). Refer to Figure 1-7.
2. Remove the yellow shipping bracket according to the following steps. Refer to Figure 1-8.
   A. Remove the screw at the top of the air vent and remove the air vent.
   B. Loosen the screw that secures the yellow shipping bracket onto the back of the compressor.
   C. Remove the bracket and then tighten the screw.
   D. Replace the air vent and secure it using the screw that was removed.
   E. Leave the yellow shipping bracket off and store it in a safe place so that if it is needed for future shipping, it is available.
3. Plug the power cord into the back of the unit and the other end into an electrical outlet that has the correct input power parameters as indicated in the General Specifications section of this manual.

**Note:** Leave the On/Off switch to the OFF position and do not apply power to the unit at this time.
Pneumatonometer Installation (continued)

4. Remove the probe from its container and place the clean tip and membrane assembly that was sent with the unit onto the probe, if one is not already installed. If a tip is not installed, perform the following:
   A. Perform the steps of the Disinfection of the Tip and Membrane Assembly as indicated in the Reprocessing Guidelines section of this manual
   B. Install the tip and membrane onto the probe tip so that the tubing is positioned approximately half way down the metal tube as shown in Figure 1-9.

Note: The correct installation of the tip and membrane assembly is that the tubing is installed approximately half way down the metal tube.

5. Plug the other end of the probe tubing into the front panel connector as follows: (Refer to Figure 1-10)
   A. Push in the probe connector into its mating connector.
   B. Ensure the connection is tight and that there is no sound of leaking air from around the connector.

6. Loosely coil the probe tubing around the probe tray using the cutouts at either end of the tray and place the probe in the tray so that the tip is not in contact with any other item.

7. If the unit includes the optional footswitch, connect the footswitch cord into the receptacle on the back panel. Refer to Figure 1-5 for the location of the footswitch receptacle.

8. Make sure that the printer has the correct thermal paper installed properly. If the unit needs thermal paper installed, refer to the Maintenance section of this manual for the installation instructions.
Application of Input Power

This section describes the Model 30 Pneumatonometer initialization.

1. Turn the instrument ON by pressing down the “I” on the front panel ON /OFF switch. At start-up, the Model 30 Pneumatonometer LCD readout will display the initial screen for several seconds as the instrument conducts a self-check test. After initialization, the unit will display the Main Menu Screen. Refer to Figure 2-1.

2. Touch the SETUP icon to enter the setup mode.

3. Touch the OK icon to set the options in the General Setup mode. Refer to Figures 2-2 and 2-3.
   A. Chart – this option sets the Chart function ON or OFF. Default is ON.
   B. Footswitch - this option sets the footswitch ON or OFF. Default is OFF.
   C. Volume – sets the volume level for the audible prompts.
   D. Brightness – sets the brightness intensity level for the LCD Screen.

4. Touch the RETURN icon to return to the Main Menu screen.
Instrument Setup (continued)

Application of Input Power (continued)

5. Touch the DOWN ARROW icon to highlight the Printout Setup option.
6. Touch the OK icon to select the Printout Setup menu. Refer to Figure 2-4.
7. Touch the UP or DOWN icons to highlight the option that needs to be changed and touch the OK icon to select the option. Refer to Figure 2-5.
8. Touch the LEFT, RIGHT, “+” or “−” icons to change the option to the appropriate setting. Refer to Figure 2-6.
9. Touch the OK icon to activate the Printout Setup menu screen.
10. If all the options are set to the desired settings, touch the RETURN icon to display to the Setup Menu screen.
11. Touch the RETURN icon to return to the Main Menu screen.

Disconnection of Input Power

1. At any time, the power switch can be set to OFF. The unit does not have a power down sequence. To terminate operation of this instrument, press the ON / OFF switch to the OFF position (O).
2. If this instrument is intended to be OFF for an extended period of time, it can be disconnected from power by detaching the power cord from the its receptacle.
Quick Verification Check

Verify the operation of the probe and base unit using the calibration verifier in the Manual mode. Refer to Figure 2-7.

1. Fill the tube of the Calibration Verifier up to the line marked 15 mmHg with filtered water.
2. From the Main Menu, touch the Manual IOP icon. A menu screen will be displayed describing a summary of the Manual IOP measurement process. Touch the OK icon to continue to the Manual IOP screen.
3. The Model 30 Pneumatonometer will display the OD and OS menu. Touch the OD or the OS icon and touch the probe of the unit to the membrane and align the tip of the probe so that white part of the probe handle is between the red and black lines on the metal tube. When the probe is between the red and black lines, the Model 30 Pneumatonometer will be ready to display the average IOP readings and its standard deviation. Refer to Figure 2-8.

Note: If the unit makes a loud vibration, remove the shipping bracket located on the back of the unit.

Note: The tone will settle to a lower tone when the probe is aligned properly. During an actual test, the tone will change to a noticeably lower pitch when the standard deviation is below 1.0 mmHg for at least three seconds. A lower tone change signifies that the instrument has acquired usable data.

After the reading is acquired, the probe can be removed from the Calibration Verifier.

Note: If the IOP is not 15 mmHg (± 2.0 mmHg), go to the Troubleshooting section of this manual for assistance. If the Troubleshooting section of the manual did not help, contact Reichert at the address or phone number in the Introduction section of this manual.

4. Touch the HOME icon to display the Main Menu.
Operating principle

The Model 30 Pneumatonometer measures intraocular pressure (IOP) non-invasively through applanation tonometry. The Pneumatonometer probe contains a gentle, floating pneumatic sensor that touches the surface of the anesthetized cornea with the exact amount of applanating force required to take a tonometry or tonography measurement. Refer to Figure 3-1.

The sensing element is a lightweight plastic tip covered with a thin, highly elastic silicone membrane. The tip is mounted on a floating piston supported by a porous bearing.

A precisely regulated flow of filtered air enters the piston from the Model 30 Pneumatonometer and travels through the end of the sensor tip until it is blocked by the membrane. When nothing is touching the membrane, air flows to the periphery of the tip, where it escapes through venting ports. However, when the tip touches the eye, the pressure against the membrane causes it to seal the vents, blocking the escape of air and building up pressure in the system. The pressure increases until it matches the IOP and stops when the eye is applanated. At this point, the membrane can no longer maintain the seal. Any increased back pressure in the system is released through the venting ports. Applanation is then automatically maintained by the pneumatic feedback system. Refer to Figure 3-2.

The Model 30 Pneumatonometer monitors the back pressure in the sensor and displays a real-time moving average of the IOP as well as its standard deviation. The Model 30 Pneumatonometer generates an audible tone to tell you when the most accurate reading is obtained. The tone will change to a noticeably lower pitch when the standard deviation among IOP readings remains below 1.0 mm Hg for three seconds. The reading remains on the LCD readout when the probe is removed from the eye.

The Model 30 Pneumatonometer in the tonography mode calculates and displays the C value (aqueous outflow coefficient) using either a two-minute or a four-minute tonography examination.
Instructions for Use

This instrument is an easy to use instrument which provides fast and accurate tonometry and optional tonography functions. Utilizing a pneumatic pump and 5mm soft silicone contact area, the Model 30 Pneumatonometer records 40 readings per second and displays real time readings of intraocular pressure on an integral graphic printer. In pulse tonometry mode, the ocular pulse waveform is charted and recorded along with IOP. There are three different tests that can be performed with the Model 30 Pneumatonometer. They are:

- **Manual Tonometry**: Displays real time readings of intraocular pressure.
- **Pulse Tonometry**: Displays real time readings of the ocular pulse waveform with a chart output.
- **Tonography**: Measurement of intraocular pressure over a two or four minute period, used to determine the rate of aqueous outflow.

Before taking measurements, there are certain conditions that should be analyzed. They are:

- **Patient preparation**.
- **General Measurement Information**.

## Patient Preparation

For manual and pulse tonometry, the patient can be either seated or supine. For tonography, the IOP must be measured first with the patient seated and then again with the patient supine. The actual tonography test is conducted with the patient in the supine position.

Use the following procedures to prepare the patient:

1. Depending on the test, have the patient seated straight in a chair, or supine.
2. Anesthetize the patient’s eye(s) according to the physician’s protocol.

**Note**: During the tests, it is best to have the patient concentrate on a fixation point to prevent the eye from wandering and causing erratic measurements.

3. Arrange the patient so that the operator is in a position to observe that the sensor membrane of the probe tip is parallel and centered on the patient’s cornea during measurements and that the probe is aligned between the red and black lines.
4. Describe the measurement procedure to the patient and what they should experience when a measurement is taken. Advise the patient to remain very still during the procedure.
Instructions for Use (continued)

Measurement Protocol

CAUTION: THE TIP AND MEMBRANE ASSEMBLY MUST ALWAYS BE CLEANED BEFORE AND AFTER MEASUREMENTS. REFER TO THE REPROCESSING SECTION OF THIS MANUAL FOR THE SUGGESTED CLEANING INSTRUCTIONS.

1. From the main menu screen, select the Manual, Pulse, or Tonography test as necessary. Refer to Figure 4-1.
2. After selecting the test, the screen will display a message to apply the probe to the patient's cornea. When touching the cornea, perform the following steps: (Refer to Figure 4-2, Measurements)

Note: The footswitch is used to start and stop the tonography test.

A. Hold the sensor lightly between the thumb and the index finger and rest the other three fingers on the patient's cheek.
B. With your other hand, lift the patient's upper eyelid.

Note: Hold the eyelid gently, excessive force applied to the eye will cause variability and inaccurate measurements.

C. Instruct the patient to focus on a fixation point.
D. Move the sensor into position along his/her line of sight and raise the back of the sensor until the piston extends directly toward the cornea.
E. Gently move the sensor toward the eye, aligning it so that the center of the tip will make initial contact close to the corneal apex.

Note: Correct alignment is required during contact to obtain valid readings.

F. As the membrane touches the cornea, continue to move the sensor handle toward the eye until the black line on the probe piston is just hidden under the white housing. Refer to Figure 4-3.

Note: Do not move the probe too far so that the red colored line on the piston is hidden under the white housing.

3. The instrument provides a series of tones when acquiring measurements (provided the volume is set high enough in the Setup mode). When a low tone is audible, the measurement data is considered usable.

Note: The tone pitch will drop noticeably when the standard deviation of the readings is below 1.0 mmHg for three seconds.
Instructions for Use (continued)

Measurements

Manual Tonometry

To measure IOP using the manual tonometry mode, follow this procedure:

**Note:** If a graphical output is desired, go to the Setup menu and set the printer to ON.

1. Touch the MANUAL IOP icon and then touch the OS or the OD icon. Refer to Figures 5-1 and 5-2.
2. Gently apply the probe to the cornea using the procedures discussed in the Measurement Protocol section. Refer to Figure 5-3.

**Note:** The pneumatonometer will continue taking readings as long as the probe maintains proper contact with the cornea. When the standard deviation between readings goes below 1.0 mmHg for three seconds, the Model 30 Pneumatonometer will lower the tone pitch indicating that usable measurements have been attained.

3. If the readings are acceptable, touch the OK icon.

**Note:** To reject all results, touch the Home icon and the system will return to the Main Menu so the test can be repeated.

4. Touch the OS or the OD icon and test the other eye.
5. Touch the PRINT icon to print the results of the test, if desired.
6. Touch the HOME icon to display the Main Menu.
**Measurements** (continued)

**Pulse Tonometry**
To measure IOP using the pulse tonometry mode, follow this procedure:

**Note:** If a graphical output is desired, go to the Setup menu and set the printer to ON.

1. Touch the PULSE IOP icon and then the OS or the OD icon. Refer to Figures 6-1 and 6-2.
2. Gently apply the probe to the cornea using the procedures discussed in the Measurement Protocol paragraph in this section. Refer to Figure 6-3.

**Note:** The tone will change when the Model 30 Pneumatonometer has sensed five ocular pulses indicating that the instrument has enough samples to compute an average. The instrument will end the test after it has sensed ten ocular pulses. The reading displayed is the average of all detected pulses.

3. If the reading is acceptable, touch the OK icon.

**Note:** To reject all results, touch the HOME icon and the system will return to the Main Menu so the test can be repeated.

4. Touch the OS or the OD icon for testing of the other eye.
5. Touch the PRINT icon to print the results of the test, if desired.
6. Touch the HOME icon to display the Main Menu.
Measurements (continued)

Tonography

The tonography test measures aqueous humor outflow. In this test, the Model 30 Pneumatonometer records changes in IOP when a weighted probe is held on the cornea for a specified time. The pneumatonometer enables the operator to conduct a two-minute or four-minute test for each eye. At the end of a test, the pneumatonometer prints a graph and tables showing the results. To measure IOP using the pulse tonometry mode, follow this procedure:

- Measure IOP with patient seated, or enter the results from a previous measurement.
- Measure IOP with the patient supine, or enter the results from a previous reading.
- Measure aqueous outflow with the patient supine using the weighted probe.
- Edit the data if necessary (Adjust Endpoints).
- Print the results (Print Statistics).

Note: The tonography option requires the Tonography Kit, P/N 16033.

CAUTION: THE TIP AND MEMBRANE ASSEMBLY MUST ALWAYS BE CLEANED BEFORE AND AFTER MEASUREMENTS. REFER TO THE REPROCESSING SECTION OF THIS MANUAL FOR THE SUGGESTED CLEANING INSTRUCTIONS.

1. With the main menu displayed, touch the TONOGRAPHY icon. Refer to Figure 7-1.
2. Touch the 2-MIN or 4-MIN icon to select the test time. Refer to Figure 7-2.
3. Touch the OD or the OS icon to select either the right eye or left eye, respectively. Refer to Figure 7-3.

Note: After the OD or the OS icon is selected, the Seated data screen is active.

Note: The tone will settle to a lower tone when the probe is aligned properly. During an actual test, the tone will change to a noticeably lower pitch when the standard deviation is below 1.0 mmHg for at least three seconds. This tone change signifies that the instrument has acquired usable data.
Instructions for Use (continued)

Measurements (continued)

Tonography (continued)

Seated IOP
1. Either touch the MEASURE or the ENTER DATA icon to acquire the Seated IOP data. Refer to Figure 7-4.
   • If the MEASURE icon is selected, then gently apply the probe to the patient’s cornea as indicated in the Measurement Protocol paragraph in this section to acquire the data.
   • If the ENTER DATA icon is selected, then touch the “+” or “–” icons as needed to enter the IOP data. Refer to Figure 7-5.
2. Touch the OK icon to enter the Seated IOP data.

Note: After the Seated IOP data is entered, the Supine data screen is active.

Supine IOP
1. Either touch the MEASURE or the ENTER DATA icon to acquire the Supine IOP data. Refer to Figure 7-6.
   • If the ENTER DATA icon is selected, touch the “+” or “–” icons as needed to enter the IOP data. Refer to Figure 7-7.
   • If the MEASURE icon is selected, then gently apply the probe to the patient’s cornea as indicated in the Measurement Protocol paragraph in this section to acquire the data.
2. Touch the OK icon to enter the supine IOP data.

Note: After the Supine IOP data is entered, the Tonography screen is active.

Note: C is the aqueous humor outflow coefficient. C is derived from the supine tonometry reading, the change in pressure from initial to final IOP, the change in volume due to pressure and the change in volume due to deformation of the cornea.

Note: P0/C is the pressure in an undisturbed eye divided by the outflow coefficient.

Note: If the probe loses contact with the eye before the end of the test, the Model 30 Pneumatonometer will abort the test. The instrument will then give a prompt to choose between START TONOGRAPHY and CALCULATOR MODE, that is, start the tonography portion of the test again or enter values manually from the measurements.
Instructions for Use (continued)

Measurements (continued)

Tonography (continued)

Supine IOP (continued)
3. Touch either the Calculator or the Tonography icon to acquire the tonography data. Refer to Figure 7-8.
   • If the Tonography icon was selected, gently apply the probe with the 10 gram weight to the patient's cornea and acquire the data until the tonography test ends. Maintain the probe alignment between the red and black lines during this test. Refer to Figure 7-10.
   • If the Calculator icon was selected, adjust the initial and final IOP values as prompted. Refer to Figures 7-11 through 7-13.
4. Touch the OK icon to enter the Tonography IOP data.
5. After all the data is acquired, touch the OK icon and then select the PRINT STATISTICS or VIEW STATISTICS option, as desired. Refer to Figure 7-9.
6. Select the END TEST option.
7. Touch the OD or OS icon to select the other eye and repeat the Tonography test as indicated above, if necessary.

Figure 7-8, Tonography

Figure 7-9, Calculator IOPs

Figure 7-10, Tonography 10g. Weight

Figure 7-11, Initial IOPs

Figure 7-12, Final IOPs

Figure 7-13, Calculator Final
Cleaning & Maintenance

Tip and Membrane Assembly

The tip and membrane assembly contacts the surface of the patient’s eye and must be cleaned to a high level of disinfection after each use to prevent cross contamination between patients. It is recommended by the manufacturer of the Model 30 Pneumatonometer to utilize the following process to achieve a high level of disinfection on the tip and membrane assembly.

**WARNING:** THE TIP AND MEMBRANE ASSEMBLY MUST BE CLEANED BEFORE AND AFTER MEASUREMENTS WITH 70% ISOPROPYL ALCOHOL FOLLOWED BY 3% HYDROGEN PEROXIDE. TO PREVENT DAMAGE TO THE EYE, THE ALCOHOL AND HYDROGEN PEROXIDE MUST BE COMPLETELY RINSED AND DRY PRIOR TO USE.

**WARNING:** DO NOT FLASH AUTOCLAVE THE TIP AND MEMBRANE ASSEMBLY. FLASH AUTOCLAVING CAN MELT THIS DEVICE.

**WARNING:** DO NOT PLACE THE PROBE, WITH A HIGH LEVEL DISINFECTED TIP, ON THE FRONT TRAY OF THE MODEL 30 PNEUMATONOMETER CONSOLE SINCE IT MAY COMPROMISE THE INTEGRITY OF THE DISINFECTION. THE TRAY SHOULD ONLY BE USED AS A STORAGE LOCATION FOR THE PROBE WHEN IT IS NOT IN USE AND PRIOR TO DISINFECTION.

**CAUTION:** DO NOT USE A WIRE BRUSH ON THE TIP AND MEMBRANE ASSEMBLY AS IT MAY DAMAGE THE MEMBRANE.

Cleaning & Disinfecting the Tip and Membrane Assembly

1. Remove the tip and membrane assembly from the probe.
2. Separate the silicone membrane from the tip.
3. **Cleaning** - Soak tip and membrane separately in 50-100 ml of 70% isopropyl alcohol for 5 minutes. To dislodge organic matter, either swirl the tip and membrane in the alcohol solution or place the alcohol container with tip and membrane into an ultrasonic cleaner.
4. Make sure there is no dust or lint in the tip air vents prior to continuing with the disinfecting process. If debris is apparent, repeat step 3.
5. **Disinfecting** - Soak the tip separately from the membrane for two 15 minute cycles in a minimum of 150-200 ml fresh 3% Hydrogen Peroxide.
6. Thoroughly rinse both tip and membrane with sterile water.
7. Allow tip and membrane to completely air dry (25-30 minutes) prior to use.
8. Replace the membrane onto the tip. The membrane should float freely on the tip. To test this, gently turn the membrane while holding the tip stem. If the membrane will not move freely on the tip, additional drying time may be necessary. If dry, inspect the tip and membrane for damage.

Inspection of the Tip and Membrane Assembly

Before and/or after reprocessing, inspect the tip and membrane for damage and wear. A slit lamp or low powered microscope can be used, as necessary. The opening in the center of the tip should be a distinct edge with no nicks or distortions. Discard the tip and membrane assembly if you see drying, cracking, deformation, or deterioration of the components.
Cleaning & Maintenance (continued)

Fuses

Fuses are located next to the power inlet. Replace fuses with only a rating as indicated on the power inlet panel. Refer to the General Specifications section in this manual. Refer to Figure 8-1.

**WARNING:** DISCONNECT POWER BEFORE ATTEMPTING TO REMOVE THE FUSES OR SERIOUS INJURY OR DEATH MAY OCCUR.

Replace the fuses in the Power Input Module with the fuses indicated in the Specifications section of this manual.

1. Remove input power to the instrument and press down on the tab in the middle of the Power Input Module to release the Fuse Holder. Refer to Figure 8-1.
2. Pull the fuse holder out of the input module.
3. Install new fuses that are indicated in the Specification section of this manual into the Fuse Holder.
4. Push the Fuse Holder into the Power Input Module until it snaps into place.

**Note:** Replacement of this fuse must be performed by qualified service personnel only.

Calibration Verifier Cornea Replacement

The artificial cornea for the Calibration Verifier can be replaced if it is damaged by replacing the cornea assembly. The cornea assembly is held in place by a o-ring. To replace the cornea assembly, perform the following steps. Refer to Figure 8-2.

1. Remove any water from the calibration verifier by pouring it out from the top of the tube until all the water is removed.
2. Place a medium flat-blade screwdriver into the slot on one side of the cornea assembly as indicated in Figure 8-2 and push one side of the cornea assembly away from the acrylic base.
3. Repeat the above step for the slot on the opposite side to remove the assembly from the acrylic base.
4. Replace with a new cornea assembly by pressing the cornea assembly into the acrylic base until it is firmly in place.

**External Cleaning**

Clean the external surfaces of this instrument at least 3 times per year using a clean, soft cloth moistened with a mild detergent solution (1 cc of liquid dish soap to one liter of clean, filtered water (filtered below 5 microns)).

Wipe down the console and front tray with disinfectant solution or mild detergent and water. Wring excess solution from cloth prior to cleaning. Moisture inside the console could cause damage. After cleaning, thoroughly dry the console with a clean, non-abrasive cloth.
Printer Paper

Install a new roll of thermal printer paper (Reichert P/N is listed in the Accessories section of this manual) as indicated. Refer to the illustration as needed.

1. Pull forward on the printer door handle to open the door. Refer to Figure 8-3.
2. Install the paper onto the paper spindle so that it unwinds counterclockwise off of the roll (the paper rolls off between the roll of paper and the printer door).
3. Feed the paper around the outside of the door and then close the printer door.
4. Align the thermal printer paper into the paper slot as shown in the adjacent picture and press the Paper Feed button on the Main Menu screen until the paper advances in and then out of the printer. Refer to Figure 8-4.

Filter Replacement

There are two filters (order Product No. 232346) in the Model 30 Pneumatonometer which should be changed semi-annually using the following procedure for each:

Compressor Filter Replacement

1. Remove the filter Access Door in the rear panel by removing the screw at the top of the door. Refer to Figure 8-5.
2. Remove the screw and filter cover from the end of the pump. Refer to Figure 8-6.
3. Remove the foam Pump Filter and replace it with the new filter from the Reichert Filter Kit (refer to the Accessories section of this manual for the part number). Refer to Figure 8-6.
4. Install the removed filter cover and screw and then tighten the screw.
5. Install the removed Access Door and screw and then tighten the screw. Refer to Figure 8-5.

Inline Filter Replacement

CAUTION: DO NOT USE EXCESSIVE FORCE WHEN REMOVING THE FILTER. IF THE INTERNAL TUBING DISCONNECTS FROM THE INLINE FILTER, THE UNIT WILL MALFUNCTION.

1. Remove the filter Access Door in the rear panel by removing the screw at the top of the door. Refer to Figure 8-5.
2. Remove the Inline Filter from by turning the connectors a quarter turn counterclockwise.

CAUTION: TAKE CARE TO AVOID KINKING THE TUBING WHEN INSTALLING THE NEW FILTER INTO THE ACCESS CAVITY.

3. Replace the filter with the new filter from the Reichert Filter Kit (refer to the Accessories section of this manual for the P/N). Refer to Figure 8-6.
4. Install the removed Access Door and screw and then tighten the screw. Refer to Figure 8-5.
# Troubleshooting

The following chart provides details of common problems and solutions for the Model 30 Pneumatonometer.

<table>
<thead>
<tr>
<th>Definition</th>
<th>Probable Cause</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screen is blank.</td>
<td>ON/OFF Switch set to OFF.</td>
<td>Press the “—” on the ON/OFF Switch.</td>
</tr>
<tr>
<td></td>
<td>Fuse(s) Blown.</td>
<td>Replace blown fuse(s).</td>
</tr>
<tr>
<td>Instrument not responding to icon touch.</td>
<td>Instrument is “locked up.”</td>
<td>Press the ON/OFF button to OFF, wait ten seconds then push it to ON.</td>
</tr>
<tr>
<td>Instrument making a loud vibrating noise.</td>
<td>Shipping bracket still attached to the unit.</td>
<td>Remove the shipping bracket as indicated in the Instrument Setup section of this manual.</td>
</tr>
<tr>
<td>Will not take a reading.</td>
<td>Patient not holding still.</td>
<td>Encourage patient to remain still.</td>
</tr>
<tr>
<td></td>
<td>Operator not positioning probe between the red and black lines.</td>
<td>Have operator refer to the Instrument Operation section of this manual.</td>
</tr>
<tr>
<td></td>
<td>Operator not holding probe perpendicular to the cornea.</td>
<td>Have operator refer to the Instrument Operation section of this manual.</td>
</tr>
<tr>
<td></td>
<td>Patient has dry eye.</td>
<td>Have patient blink eyes.</td>
</tr>
<tr>
<td></td>
<td>Unit needs reboot of hardware.</td>
<td>Unplug unit, wait 2 minutes then apply input power.</td>
</tr>
<tr>
<td>Readings are low.</td>
<td>Air leak in the system.</td>
<td>Check hoses for air leaks.</td>
</tr>
<tr>
<td></td>
<td>Probe connector not fully engaged on unit.</td>
<td>Check probe connection on unit.</td>
</tr>
<tr>
<td></td>
<td>Filter is becoming clogged.</td>
<td>Change Filter - Refer to Filter Kit P/N in the Accessory section.</td>
</tr>
<tr>
<td>Readings are high.</td>
<td>Dirt/Contamination in the system.</td>
<td>Check hoses for dirt or contamination.</td>
</tr>
<tr>
<td></td>
<td>Tip and membrane not installed properly.</td>
<td>Refer to the Tip Installation Procedure in this manual.</td>
</tr>
<tr>
<td></td>
<td>Filter is becoming clogged.</td>
<td>Change Filter - Refer to Filter Kit P/N in the Accessory section.</td>
</tr>
<tr>
<td>Screen difficult to see.</td>
<td>Contrast is set too low.</td>
<td>Adjust contrast in Setup menu.</td>
</tr>
<tr>
<td>Printer not printing.</td>
<td>Printer out of paper.</td>
<td>Replace paper with Reichert thermal printer paper as indicated in the Accessories section of this manual.</td>
</tr>
<tr>
<td></td>
<td>Printer paper in backwards.</td>
<td>Reverse the thermal printer paper.</td>
</tr>
<tr>
<td></td>
<td>Not using thermal paper from Reichert.</td>
<td>Replace paper with Reichert thermal printer paper as indicated in the Accessories section of this manual.</td>
</tr>
<tr>
<td>Calibration verifier cornea damaged</td>
<td>Replace the Cornea Assembly.</td>
<td>Replace to the Maintenance Section of this manual.</td>
</tr>
</tbody>
</table>
# Specifications

This section contains the specifications for the Model 30 Pneumatonometer.

<table>
<thead>
<tr>
<th><strong>Physical Dimensions</strong></th>
<th><strong>Instrument Console</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Size:</td>
<td>Weight, unpacked: 10.0 lbs. (4.54 Kg)</td>
</tr>
<tr>
<td>Height:                5.25 in. (13.3 cm)</td>
<td></td>
</tr>
<tr>
<td>Width:                 14.0 in. (35.6 cm)</td>
<td></td>
</tr>
<tr>
<td>Depth:                 10.5 in. (26.7 cm)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Instrument Probe</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Size:</td>
</tr>
<tr>
<td>Outside Diameter:   0.5 in. (1.3 cm)</td>
</tr>
<tr>
<td>Length:              4.25 in. (10.8 cm)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Tonometry</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>IOP Range: 5-80 millimeters of mercury (mmHg)</td>
</tr>
<tr>
<td>Measurement Accuracy (at 95% confidence)</td>
</tr>
<tr>
<td>0 - 40 mmHg ± 1.5 mmHg</td>
</tr>
<tr>
<td>40 - 80 mmHg ± 3.5 mmHg</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Tonography</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Effective Sample Rate: 40 hertz</td>
</tr>
<tr>
<td>Real-time Chart Printing Speed: 50 millimeters per minute</td>
</tr>
<tr>
<td>Chart Scale: 0 to 80 millimeters of mercury (mmHg)</td>
</tr>
<tr>
<td>Test Duration: 2 or 4 minutes</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Electrical</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Part Number (P/N): P/N 16030 P/N 16031 P/N 16032</td>
</tr>
<tr>
<td>Voltage: 120 volts AC 50/60 Hz 230 volts AC 50/60 Hz 100 volts AC 50/60 Hz</td>
</tr>
<tr>
<td>Current: 2.0 A 1.6 A 2.0 A</td>
</tr>
<tr>
<td>Watts: 43 VA 50 VA 57 VA</td>
</tr>
<tr>
<td>Fuses: Time-Lag (T 2.0 A L 250V), 5 x 20mm, RoHS Time-Lag (T 1.6 A L 250V), 5 x 20mm, RoHS Time-Lag (T 2.0 A L 250V), 5 x 20mm, RoHS</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Operational Conditions</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Environmental:</td>
</tr>
<tr>
<td>The environmental conditions are as follows:</td>
</tr>
<tr>
<td>Operating:</td>
</tr>
<tr>
<td>Temperature: 10°C (50°F) to 35°C (95°F)</td>
</tr>
<tr>
<td>Relative Humidity: 30% to 75% (non-condensing)</td>
</tr>
<tr>
<td>Atmospheric Pressure: 80 (23.6 in. Hg) to 106 kPa (31.3 in. Hg)</td>
</tr>
<tr>
<td>Transportation &amp; Storage:</td>
</tr>
<tr>
<td>Temperature: -20°C (-4°F) to +70°C (158°F)</td>
</tr>
<tr>
<td>Relative Humidity: 10% to 80% (non-condensing)</td>
</tr>
<tr>
<td>Atmospheric Pressure: 70 (20.7 in. Hg) to 106 kPa (31.3 in. Hg)</td>
</tr>
<tr>
<td>Exposure to extreme temperature conditions indicated above must not exceed 15 weeks.</td>
</tr>
</tbody>
</table>
Disposal
This product does not generate any environmentally hazardous residues. At the end of its product life, follow your local laws and ordinances regarding the proper disposal of this equipment.

Software Revision
The software revision can be obtained by contacting Reichert, Inc. The serial number identifies the manufacture date and will provide access to the software version.

Classifications
The Model 30 Pneumatonometer is classified as Class I Equipment.

Class I Equipment is equipment in which protection against electric shock does not rely on basic insulation only, but which includes an additional safety precaution in that means are provided for the connection of the equipment to a protective earth conductor in the fixed wiring of the installation in such a way which accessible metal parts cannot become live in the event of a failure of the basic insulation.

The Model 30 Pneumatonometer is classified as Type B Equipment.

Type B Equipment provides an adequate degree of protection against electrical shock, particularly regarding allowable leakage currents and reliability of the protective earth connection.

The Model 30 Pneumatonometer is classified as IPX0 Equipment.

IPX0 Equipment is ordinary equipment enclosed without protection against ingress of water.

According to the mode of operation, the Model 30 Pneumatonometer is a Continuous Operation instrument.
Guidance & Manufacturer’s Declarations

Table 201 – Guidance and Manufacturer’s Declaration

Electromagnetic Emissions
All Equipment and Systems

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Compliance</th>
<th>Electromagnetic Environment - Guidance -</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF Emissions CISPR 11</td>
<td>Group 1</td>
<td>The Model 30 uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td></td>
<td>Class A (16030)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Class B (16031)</td>
<td></td>
</tr>
<tr>
<td>Harmonics IEC 61000-3-2</td>
<td>Class A</td>
<td>The Model 30 is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies building for domestic power.</td>
</tr>
<tr>
<td>Flicker IEC 61000-3-3</td>
<td>Complies or N/A</td>
<td></td>
</tr>
</tbody>
</table>

**Note:** The emissions characteristics of model 16030 make it suitable for use in industrial areas and hospitals (CISPR 11 class A). If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radio-frequency communication services. The user might need to take mitigation measures, such as relocating or re-orienting the equipment.
## Electromagnetic Immunity

The Model 30 is suitable for use in electromagnetic environment specified below. The customer or user of the Model 30 should ensure 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>ESD IEC 61000-4-2</td>
<td>±2kV, ±4kV, ±6kV, ±8kV Contact ±2kV, ±4kV, ±8kV, ±15kV Air</td>
<td>±2kV, ±4kV, ±6kV, ±8kV Contact ±2kV, ±4kV, ±8kV, ±15kV Air</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are synthetic, the R/H should be at least 30%.</td>
</tr>
<tr>
<td>EFT IEC 61000-4-4</td>
<td>±0.5kV, ±1kV, ±2kV Mains ±0.5kV, ±1kV I/Os</td>
<td>±0.5kV, ±1kV, ±2kV Mains ±0.5kV, ±1kV I/Os</td>
<td>Mains power quality should be that of a typical residential, commercial or hospital environment.</td>
</tr>
<tr>
<td>Surge IEC 61000-4-5</td>
<td>±0.5kV, ±1kV Differential ±0.5kV, ±1kV, ±2kV Common</td>
<td>±0.5kV, ±1kV Differential ±0.5kV, ±1kV, ±2kV Common</td>
<td>Mains power quality should be that of a typical residential, commercial or hospital environment.</td>
</tr>
<tr>
<td>Voltage Dips/Dropout IEC 61000-4-11</td>
<td>&gt;95% Dip for 0.5 Cycle 60% Dip for 5 Cycles 30% Dip for 25 Cycles &gt;95% Dip for 5 Seconds</td>
<td>&gt;95% Dip for 0.5 Cycle 60% Dip for 5 Cycles 30% Dip for 25 Cycles &gt;95% Dip for 5 Seconds</td>
<td>Mains power quality should be that of a typical residential, commercial or hospital environment. If the user of the Model 30 requires continued operation during power mains interruptions, it is recommended that the Model 30 be powered from an uninterruptible power supply or battery.</td>
</tr>
<tr>
<td>Power Frequency 50/60Hz Magnetic Field IEC 61000-4-8</td>
<td>30A/m</td>
<td>30A/m</td>
<td>Power frequency magnetic fields should be that of a typical residential, commercial or hospital environment.</td>
</tr>
</tbody>
</table>
## Guidance and Manufacturer’s Declaration – Electromagnetic Immunity

The Model 30 is intended for use in the electromagnetic environment specified below. The customer or user of the Model 30 should ensure 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>(V1) = 3 Vrms</td>
<td>Portable and mobile RF communications equipment should be no closer to any part of the Model 30, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
<tr>
<td></td>
<td>3 Vrms</td>
<td>(V1) = 6 Vrms in ISM bands</td>
<td></td>
</tr>
<tr>
<td></td>
<td>150 kHz to 80 MHz</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>6 Vrms</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>150 kHz to 80 MHz in ISM bands</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Radiated RF</td>
<td>IEC 61000-4-3</td>
<td>(E1) = 3 V/m</td>
<td>Recommended Separation Distance:</td>
</tr>
<tr>
<td></td>
<td>80 MHz to 2.7 GHz @ 3 V/m</td>
<td></td>
<td>d=(3.5/V1)(Sqrt P)</td>
</tr>
<tr>
<td></td>
<td>80 MHz to 2.7 GHz @ 10 V/m</td>
<td></td>
<td>d=(3.5/E1)(Sqrt P)</td>
</tr>
<tr>
<td></td>
<td>(E1) = 10 V/m</td>
<td></td>
<td>80 to 800 MHz</td>
</tr>
<tr>
<td></td>
<td>800 MHz to 2.7 GHz</td>
<td></td>
<td>d=(7/E1)(Sqrt P)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>800 MHz to 2.7 GHz</td>
</tr>
</tbody>
</table>

Where P is the max output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m).

Field strengths from fixed transmitters, as determined by an electromagnetic site survey, should be less than the compliance levels in each frequency range.

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

---

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

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

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

* Over the frequency range 150 kHz to 80 MHz, field strengths should be less than [V1] V/m.

* The ISM (industrial, scientific and medical) bands between 0,15 MHz and 80 MHz are 6,765 MHz to 6,795 MHz; 13,553 MHz to 13,567 MHz; 26,957 MHz to 27,283 MHz; and 40,66 MHz to 40,70 MHz. The amateur radio bands between 0,15 MHz and 80 MHz are 1,8 MHz to 2,0 MHz, 3,5 MHz to 4,0 MHz, 5,3 MHz to 5,4 MHz, 7 MHz to 7,3 MHz, 10,1 MHz to 10,15 MHz, 14 MHz to 14,2 MHz, 18,07 MHz to 18,17 MHz, 21,0 MHz to 21,4 MHz, 24,89 MHz to 24,99 MHz, 28,0 MHz, to 29,7 MHz and 50,0 MHz to 54,0 MHz.
Table 206 – Recommended Separation Distances between Portable and Mobile RF Communications Equipment and the Model 30 for ME Equipment and ME Systems that are NOT Life-supporting.

Guidance and Manufacturer’s Declaration - Electromagnetic Immunity

<table>
<thead>
<tr>
<th>Max Output Power of Transmitter (W)</th>
<th>Separation (m) 150kHz to 80 MHz</th>
<th>Separation (m) 80 to 800 MHz</th>
<th>Separation (m) 800MHz to 2.7 GHz</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01</td>
<td>d=(3.5/V1)(Sqrt P)</td>
<td>0.1166</td>
<td>0.2333</td>
</tr>
<tr>
<td>0.1</td>
<td>0.3689</td>
<td>0.3689</td>
<td>0.7378</td>
</tr>
<tr>
<td>1</td>
<td>1.1666</td>
<td>1.1666</td>
<td>2.3333</td>
</tr>
<tr>
<td>10</td>
<td>3.6893</td>
<td>3.6893</td>
<td>7.3786</td>
</tr>
<tr>
<td>100</td>
<td>11.6666</td>
<td>11.6666</td>
<td>23.3333</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance (d) in meters (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 80 MHz and 800 MHz, 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.
### Guidance & Manufacturer’s Declarations (continued)

**Table 9 – Guidance and Manufacturer’s Declaration**

**Electromagnetic Immunity**

**Immunity to Proximity Fields from RF Wireless Communications Equipment**

#### Guidance and Manufacturer’s Declaration - Electronic Immunity

The Model 30 is intended for use in the electromagnetic environment as specified below related to proximity fields from RF wireless communications equipment.

<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>Test Frequency (MHz)</td>
<td>Band (MHz)</td>
<td>Service (MHz)</td>
<td>Modulation</td>
</tr>
<tr>
<td>385</td>
<td>380-390</td>
<td>TETRA 400</td>
<td>Pulse Modulation 18 Hz</td>
</tr>
<tr>
<td>450</td>
<td>430-470</td>
<td>GMR 460, FRS 460</td>
<td>FM ±5 kHz deviation 1 kHz sine</td>
</tr>
<tr>
<td>710</td>
<td>704-787</td>
<td>LTE Band 13, 17</td>
<td>Pulse Modulation 217 Hz</td>
</tr>
<tr>
<td>780</td>
<td>800-960</td>
<td>GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE Band 1, 3, 4, 25, UMTS</td>
<td>Pulse Modulation 18 Hz</td>
</tr>
<tr>
<td>810</td>
<td>1700-1990</td>
<td>GSM 1800; CDMA 1900; GSM 1900; DECT; LTE Band 1, 3, 4, 25, UMTS</td>
<td>Pulse Modulation 217 Hz</td>
</tr>
<tr>
<td>870</td>
<td>1700-1990</td>
<td>Bluetooth WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7</td>
<td>Pulse Modulation 217 Hz</td>
</tr>
<tr>
<td>930</td>
<td>5100-5800</td>
<td>WLAN 802.11 a/n</td>
<td>Pulse Modulation 217 Hz</td>
</tr>
</tbody>
</table>
Warranty

This product is warranted by Reichert Technologies ("Reichert") against defective material and workmanship under normal use for a period of one year from the date of invoice to the original purchaser. (An authorized dealer shall not be considered an original purchaser.) Under this warranty, Reichert’s sole obligation is to repair or replace the defective part or product at Reichert’s discretion.

This warranty applies to new products and does not apply to a product that has been tampered with, altered in any way, misused, damaged by accident or negligence, or that has the serial number removed, altered or effaced. Nor shall this warranty be extended to a product installed or operated in a manner not in accordance with the applicable Reichert instruction manual, nor to a product that has been sold, serviced, installed or repaired other than by a Reichert factory, Technical Service Center, or authorized Reichert Technologies Dealer.

Lamps, bulbs, charts, cards and other expendable items are not covered by this warranty.

All claims under this warranty must be in writing directed to the Reichert factory, Technical Service Center, or authorized instrument dealer making the original sale and must be accompanied by a copy of the purchaser's invoice.

This warranty is in lieu of all other warranties implied or expressed. All implied warranties of merchantability or fitness for a particular use are hereby disclaimed. No representative or other person is authorized to make any other obligations for Reichert. Reichert shall not be liable for any special, incidental, or consequent damages for any negligence, breach of warranty, strict liability or any other damages resulting from or relating to design, manufacture, sale, use or handling of the product.

PATENT WARRANTY

If notified promptly in writing of any action brought against the purchaser based on a claim that the instrument infringes a U.S. Patent, Reichert will defend such action at its expense and will pay costs and damages awarded in any such action, provided that Reichert shall have sole control of the defense of any such action with information and assistance (at Reichert’s expense) for such defense, and of all negotiation for the settlement and compromise thereof.

PRODUCT CHANGES

Reichert reserves the right to make changes in design or to make additions to or improvements in its products without obligation to add such to products previously manufactured.

CLAIMS FOR SHORTAGES

We use extreme care in selection, checking, rechecking and packing to eliminate the possibility of error. If any shipping errors are discovered:

1. Carefully go through the packing materials to be sure nothing was inadvertently overlooked when the unit was unpacked.

2. Call the dealer you purchased the product from and report the shortage. The materials are packed at the factory and none should be missing if the box has never been opened.

3. Claims should be filed within 30 days.

CLAIMS FOR DAMAGES IN TRANSIT

Our shipping responsibility ceases with the safe delivery in good condition to the transportation company. Claims for loss or damage in transit should be made promptly and directly to the transportation company.

If, upon delivery, the outside of the packing case shows evidence of rough handling or damage, the transportation company’s agent should be requested to make a “Received in Bad Order” notation on the delivery receipt. If within 48 hours of delivery, concealed damage is noted upon unpacking the shipment and no exterior evidence of rough handling is apparent, the transportation company should be requested to make out a “Bad Order” report. This procedure is necessary in order for the dealer to maintain the right of recovery from the carrier.