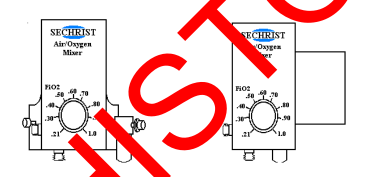
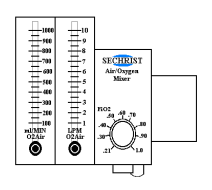


3500 / 3500HL Series Air / Oxygen Mixer USER'S MANUAL





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FEDERAL LAW (USA) RESTRICTS THIS DEVICE TO SALE BY OR ON THE ORDER OF A PHYSICIAN.

P/N 100001 Rev. 18

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INTRODUCTION

We at Sechrist Industries, Inc. thank you for choosing a Sechrist air/oxygen mixer. We also caution you that before attempting to use the mixer in a patient care setting, you must make yourself thoroughly familiar with the instructions in this manual and any product labeling. Throughout this manual, warnings, cautions, and notes will be utilized to bring your attention to particularly important matters.

USER / OWNER RESPONSIBILITY

WARNING

It is the responsibility of the procuring organization passive they have the tooling, equipment and training necessary to perform the tasks they are procuring components or kits for.

WAR

Hospitals or users who perfects certa reparand/or periodic service without undergoing training by Secretist Industry assumes sole responsibility for any malfunction, which results from improper usage, faulty maintenance, improper or unauthorized repairs, demage of teration reformed.

The Sechrist mixed will per rm in accordance with the specifications and descriptions contained within this manual and accompanying labeling when the mixer is operated and maintained / accordance with the instructions contained within this manual and other accompaning doc tion. Do not attempt to operate this equipment before reading anderst The mixer should be checked nding these instructions. periodally as specific within this manual (see Routine Maintenance section). A ective oductorald never be used in a clinical setting. Any necessary repair should be vided the Sechrist home offices in Anaheim, CA or by an individual trained and aut by Schrist Industries. However, Sechrist recognizes that some hospitals and her use, maintain their own service groups (biomedical engineers and technicians) who perform certain repairs and/or periodic service. Given this, Sechrist does provide re components and kits for such effort.

WARNINGS & CAUTIONS

WARNINGS indicate the possibility of personal injury or death to the patient and/or operator of the device.

<u>CAUTIONS</u> indicate the potential of damage to equipment and/or other property if the caution is ignored.

NOTES call attention to statements that are intended to supplement or emphasize basic instructions contained within this manual.

1

WARNING

It is the responsibility of the procuring organization to assure they have the tooling, equipment and training necessary to perform the tasks they are procuring components or kits for.

WARNING

No modification of this equipment is allowed. Do not modify this equipment without authorization from the manufacturer. If this equipment is modified, appropriate inspection and testing must be conducted to ensure continued safe use of the equipment.

WAP

Hospitals or users who perform certain very as and, periodic service without undergoing training by Sechrist Hobstries assumes sole responsibility for any malfunction, which results from improper usage faulty maintenance, improper or unauthorized repairs, damage of alteration performed.

WARNING

The user of the Sechrist Vir/Oxygen Mixer shall have sole responsibility for any malfunction, which results from improper usage, faulty maintenance, improper and/or unauthoused repair plamage or alteration performed by anyone other than Sechrist Industrie

WARNING

darm, ypass conditions must be corrected swiftly, as the selected oxygen concentration, will not be delivered during a bypass situation.

WARNING

Liquid water or other contaminants in either supply gas, particularly in the air supply, will cause malfunction of this equipment and any attached equipment. Supply gases should meet gas dryness of .0045 mg water per cubic centimeter of gas.

2

WARNING

Oxygen concentration must be monitored downstream from the mixer with a suitable, calibrated oxygen analyzer, equipped with alarms that can be set for high and low FIO₂'s. FIO₂'s should then be adjusted to maintain appropriate blood gas concentrations.

WARNING

The mixer is designed to mix air and O_2 only; do pt modify the inlets to accommodate any other source gases.

WARN NG

Oxygen vigorously accelerates combustic. To ay of prolosion hazard, do not expose the mixer to any instruments or other equipper that may have been contaminated by oil or grease. Gas supplied to the formula be extremely clean (no more than 25 parts per million (ppm) of gase us hydrocarbons callowed.) A high concentration of hydrocarbons in the gas supplies a fire hazar

WARNING

The mixer are like alarm may not function when both air and O_2 supply pressures are less than the minimum specified inlet pressure.

WARNING

To outlet have the capability of providing gas pressures equal to the inlet pressures. The ore, and attached equipment must provide safety relief protection in order to prevent pressures from being delivered to patients.

WARNING

Whenever a patient is attached to respiratory care equipment, constant attendance is required by qualified personnel. The use of alarm or monitoring systems does not provide absolute assurance of a warning for every possible system malfunction. In addition, some problems may require immediate attention.

WARNING

Excessive supply pressures (> 70 psi, 482 kPa) may result in mixer damage or malfunction. Use of a suitable supply gas regulation system is necessary.

WARNING

A Sechrist air/oxygen gas mixer is a sophisticated medical device designed for use by qualified personnel under the direction of a qualified physician.

WARNING

This product should only be maintained and repaired by a Sechrist Industries factory-trained technician or by written instructions from Sechrist Industries. This product should not be modified in any way, except with prior witten approval of Sechrist Industries. Unapproved modifications can result in death or prious injury.

WARN NG

The mixer does not contain gas-sterilizing filters are cill supply the same quality of gas supplied from the gas sources. Use of applicate gas punty and gas line filters is the responsibility of the user.

WARD NG

When the Sechrist are is used to supplement respiratory equipment, the user must refer to and follow the intructions provided by the manufacturer of the respiratory equipment.

CAUTION

Immerse the mixer in any solution. Do not sterilize.

CAUTION

This precision gas-mixing device may become nonfunctional or damaged if used whout the watertrap assembly and filters provided.

CAUTION

Before using this mixer, verify that the performance verification procedure has been performed by a qualified individual.

SUMMARY OF SYMBOLS

SYMBOL	REFERENCE	MEANING
Ţ.	ISO 7000-0434B	Caution
	BS EN ISO 15223- 1:2012	n Surer
	BS EN ISC \5223 1:2012	Date of Manufacture
Ti /	ISO 7000-1 41	The symbol indicates to user/operator to refer to instruction manual/ booklet
	ISO 7010-W001	Warning, prohibition or mandatory action

INDICATIONS FOR USE

Intended Use

The purpose of the device is to enable qualified personnel to mix medical-grade air and medical-grade oxygen, at operator selected ratios, for delivery to patients through various types of respiratory care equipment.

The Sechrist air/oxygen mixer is a precision pressure regulation and proportioning device, which is designed to accurately mix medical grade air and medical grade oxygen (O₂). The mixer can provide for FIO₂'s of .21 to 1.0 for delively to a variety of eter index safety system respiratory devices. The mixer receives air and oxygen (D.I.S.S.) inlet connections at a nominal pressure 50 psi. (344 Pa). The unit will operate satisfactorily with inlet pressures of 2 - 70 i (207 - 482 Pa) providing that the pressures are within 20 psi (138 kPa) one and her. The Sechrist air/oxygen mixers may be indicated whenever precise nce rations oxygen are required for nits appriate configuration, may be found clinical applications. Use of the mix throughout the healthcare envir ment. es in de but may not be limited to bedside delivery of precise oxy en concentra ons directly to the patient or delivery of precise FIO₂'s to other equipn nt, such as a entilator, isollettes, or resuscitation equipment.

CONTRAUNDICA YONS

While supplemental exygen therapy is not without possible side effects, such as absorption at class, and oxygen toxicity, the detrimental effects of oxygen should new prevent its use your indicated ¹.

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¹ Eagan, M. Eagan's Fundamentals of Respiratory Care, Fifth Edition 1999

OPERATING INSTRUCTIONS

- Before using the mixer, verify that performance verification has been completed.
- ♦ If applicable, connect the mixer outlet to the inlet device that will be delivering the oxygen concentration to the patient.
- ◆ From reliable, pressure regulated gas sources, connect both medical grade air and medical grade oxygen to the mixer inlets utilizing appropriate gas hoses (P/N IV 308 air supply hose and P/N IV 309 oxygen supply hose

NOTE: The alarm/bypass will activate when as the case is connected. The alarm will reset upon the connection to the stand gas supp

- ◆ Using the calibrated control knob, sect the desired oxygen concentration (FIO₂) from .21 to 1.0.
- If the configuration includes a hormeter's, initiate gas flow through the flowmeter(s) by tuning the knob(s) on the flowmeter (s) counter clockwise to the desired flow setting.
- Begin the operation of the attack envery unit, if applicable.
- After the selected gas in ture has washed out the room air from the delivery unit, analyze and monitor the clivered gas concentration with a calibrated oxygen analyser. Appropriately set the high and low alarm limits on the analyzer.
- respondically observe the watertrap assembly for the accumulation of moisture. Moreover some observed from the water trap assembly by depressing the valve at the attom of the watertrap bowl.
- Pendically observe the oxygen analyzer and evaluate the delivered FIO₂.

SPECIFICATIONS

Multiple configurations are available, with and without attached flowmeter(s). All models utilize the same gas mixer and therefore the following specifications apply to all configurations.

 FIO_2 .21 +.01 to 1.0 – 0.1

Accuracy * $\pm 3\%$

(high flow configurations) at least 100 lpm @ an FIO₂ f .60 with inlet pressures

of 50 psi (344 km. Surely range of 30-70 psi produces an output flow with a range of 70-150 lpm

(low flow configurations) at least 40 pm @ FIO₂ of .60 with inlet pressures

of 50 p. (344) Supply range of 30-70 psi produces an exput flow within a range of 29-60 lpm

Supply Pressures **

Nominal 50 psi (344 kPa)± 10 psi (68 kPa) (@ 4.0 standard

ic feet er minute (SCFM) min. flow)

Minimum 30 psi (207 kPa)

Maxin .m 70 psi (482 kPa)

Bleed Flo.

(high w co. Lations) 8.0 to 10.0 lpm @ 16 lpm flow setting

flow configurations) 2.5 to 4.5 lpm @ 8 lpm flow setting

Dimensions (without flow meters)

Height 6 inches (15.24 cm)

Width 6 inches (15.24 cm) (pole mount)

6 ½ inches (16.51 cm) (wall mount)

Depth 6 inches (15.24 cm) (pole mount)

5 ½ inches (13.97 cm) (wall mount)

Weight 6 lbs. (2.73 kg)

Service Life 2 years

Optional flowmeters

1 - 10 lpm	\pm 3% of full scale
1 – 15 lpm	\pm 3% of full scale
0 – 16 lpm	\pm 3% of full scale
2 - 20 lpm	\pm 3% of full scale
2 – 32 lpm	\pm 3% of full scale
3 - 30 lpm	± 3% of full sea.
100 - 1000 ml/min	±3% of fu scale

*NOTE: The mixer will maintain the dever a FIO_2 manin \pm 1% of the selected concentration with small fluctuation of the script pressure. The additional 2% error results from the readability of the set point and severe results.

**NOTE: The outlet pressure of the mixer all always be slightly lower than the lower of the two supply pressures. Some respiratory equipment attached to the mixer may require closer tolerators if so, constant at the manufacturer of that equipment.

***NOTE: The bleed flow located on the bottom of the proportioning module and is necessary at order to mainta. FIO₂ accuracy at very low flow settings.

Optiona Accordies

The Polywing oper or detachable inlet pressure hoses comply with Compressed s Association (CGA) V-1, V-5, and G-4.1:

```
Ref. IV 308 14 foot (4.27 m) Air Supply Hose
14 foot (4.27 m) Oxygen Supply Hose
```

NERFORMANCE VERIFICATION

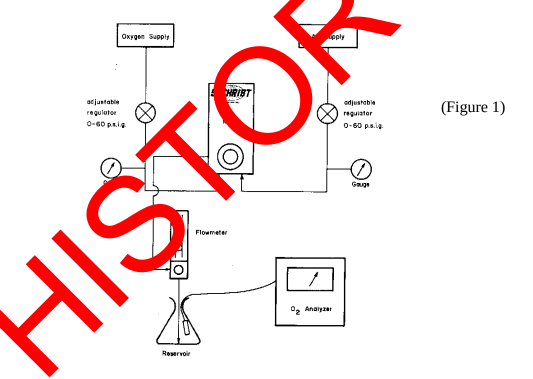
Prior to each clinical usage, the user should perform an alarm test and analyze the full FIO₂ range. With an accurately calibrated oxygen analyzer, the user should analyze the FIO₂ at the following settings; 21%, 40%, 60%, 80%, and 100%. Additionally, the user should briefly disconnect one supply gas to assure that the bypass/alarm system is functioning. With a single supply gas disconnected, the audible alarm should sound and the analyzed FIO₂ should indicate the FIO₂ of the single supply gas; i.e. 21% if the oxygen was disconnected and 100% if the air supply was disconnected.

The following more extensive procedure should be performed at least once a month, or more frequently as indicated or desired.

This procedure provides a means of determining if the mixer is functioning in accordance with the design specifications. This verification is intended to be performed in the health care setting by qualified personnel. The procedure should be followed exactly as outlined. If the mixer fails to meet the established standards, it should be removed from clinical application until calibration and/or service is accomplished (see troubleshooting section or service manual)

NOTE: It is strongly recommended that personnel responsible for performance verification testing keep accurate records of testing activities.

The performance verification process requires a simple contige tion as diagramed below.



- Connect the mixer to the supply gases with independently adjustable pressure regulators.
- Connect a flowmeter to the mixer outlet.
- Direct the flow from the flowmeter to a reservoir (e.g. a bottle or tube) making sure that no room air is being entrained to dilute the mixture.
- ❖ Place a calibrated O₂ analyzer probe within the reservoir.

Test for overall accuracy

- Set both supply pressures to 50 psi (344 kPa)
- Set the flowmeter to 8 lpm for configurations with the following flowmeters; 0-10 lpm, 1-15 lpm, 0-16 lpm, and 100-1000 ml/min.
- Set the flowmeter to 15 lpm for configurations with the following flowmeters; 2-20 lpm, 2-32 lpm, and 3-30 lpm.
- Compare the O_2 analyzer readings at the following settings. Since the mixer has an overall accuracy of \pm 3% and if the analyzer accuracy is thin \pm 1%, the following comparisons should agree within \pm 4% points.
 - **.**21
 - **.**40
 - **.**60
 - **.**80
 - **4** 1.0

Test for accuracy with ving inlet pressures.

- Set the FIO₂ to .60 with the inlet pressures at 50 psi (344 kPa).
- * Verify setting a uracy comparing the setting with the analyzed value.
- Set O₂ Sure to 40 psi (276 kPa) leaving the air supply at 60 psi (414 kPa).
- te the malyzer reading.
- ❖ Set the O₂ pressure to 60 psi (414 kPa) and the air supply to 40 psi (276 kPa)
- Note the analyzer reading.
- ❖ Analyzed O₂ concentrations should vary by no more that 2% with the above pressure changes.

Test the alarm module function.

- Set supply pressures to 50 psi (344 kPa).
- Set the FIO₂ to .60.

- Reduce the air supply pressure to 24 psi. (166 kPa).
- ❖ The audible alarm should sound within the following pressure range of 24-28 psig, and the O₂ analyzer should read 100%.
- Slowly increase the air supply pressure to 50 psi (344 kPa). The alarm should cease and reset prior to obtaining a supply pressure of 40 p.s.i. (276 kPa).
- Reduce the O₂ supply pressure to 24 psi (166 kPa).
- The audible alarm should sound within the following passure range of 24-28 psig, and the O₂ analyzer should read 21%.
- Slowly increase the O₂ supply pressure to 50 psi (344 kPa). The alarm should cease and reset prior to obtaining a supply pressure of 10 psi (276 kg).

Check the inlet filters.

- To test the flow through the through the first and air inlet filter assemblies, a small test port has been provided on the rear of the fixer just above the air inlet.
- Turn both supply goes of and allow the gases to cease flowing. With a 1/4 inch hex nut driver, remove the two from the test port and install a 10-32 threaded nipple.
- Connect occurate preserve gauge (0-60 psi) (0-414 kPa) to the nipple.
- Turn the supples on and set the flow to 16 lpm.
- the mixer FIC control to .21.
- Obserthe pressure registered by the gauge connected to the test port.
- The difference between the test gauge pressure and the supply pressure should vary by no more than 5 psi (34 kPa).
 - ❖ If the pressure differential is > 5 psi (34 kPa) replace the inlet filters as outlined in the routine maintenance section.

THEORY OF OPERATION

The Sechrist air-oxygen mixer is a precision pressure regulation and proportioning device which is designed to accurately mix medical-grade Air (air) and medical-grade Oxygen (O₂) to any selected FIO₂ between 21% and 100% for delivery to various types of respiratory care equipment. To accomplish this task, the Sechrist mixer is composed of three major components or modules. The balancing module, the proportioning module, and the alarm/bypass module.

Gas is delivered to the balancing module where inlet pressures are equalized. Supply pressures that do not meet the minimum specified pressure, may result in the device not functioning to the specification. Supply pressures that exceed the maximum specified pressure, may result in device damage or malfunction. Duckbill valves, positioned between the diaphragm and inlet filters, prevent reverse gas flow. Gas then travels to the proportioning module where the gases are mixed to the user-defined concentration. A continual flow of gas supports the alarm/bypass module, which provides an audible alarm in the event of either a significant loss of supply gas pressure or a loss of a single supply gas.

Filters are incorporated in both of the gas inlet connections. Dne-way check valves are also located in both of the gas inlet connections to prevent the cross-contamination of one gas supply to the other source.

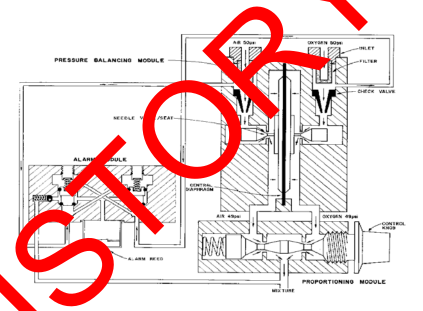


Figure 2

ROUTINE MAINTENANCE

NOTE: The manufacturer will make available on request circuit diagrams, component part lists, descriptions, calibration instructions, or other information that will assist trained service personnel to repair those parts of medical equipment that are designated by the manufacturer as repairable by service personnel.

Routine maintenance of the mixer is limited to periodic performance verification, replacement of the inlet filters and cleaning of the exterior surfaces. A mixer in need of calibration or service should not be used until the necessary procedures are performed and the equipment has been tested to determine that it is functioning properly. Calibration and servicing may only be accomplished by personnel trained and

authorized to do so by Sechrist Industries. Routine maintenance, as defined in this manual, may be performed by a competent individual having experience in the maintenance of devices of this nature. Parts designated within this manual should be replaced only with parts manufactured or sold by Sechrist Industries.

Routine maintenance procedure

Inlet filters

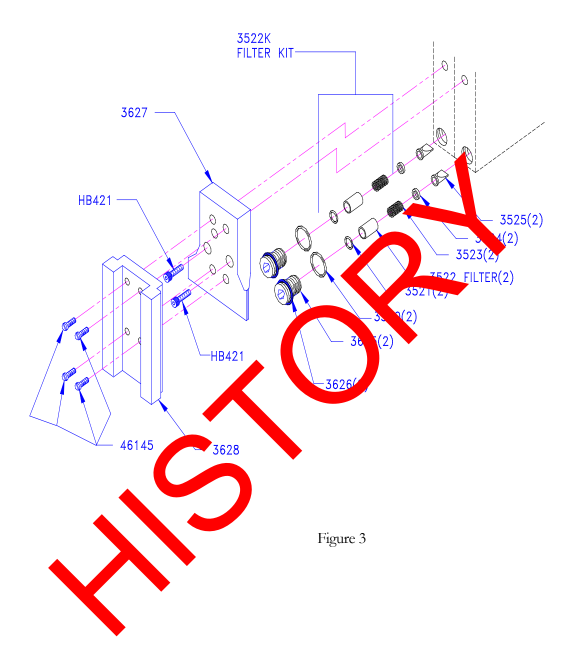
- Periodically replace the air water trap/inlet filter (PN 3529E in Figure 4) at least every 6 months or as needed.
- Replace the internal sintered stainless steel (P. N 3522K in Figure 3) at least every 6 months or as needed.

CA VION

Do not immerse the mater in any soution. Do not attempt to sterilize.

Cleaning

Exterior surfaces of the mixer may be wiped clean with a mild soap solution or a light disinfectant polution. Do not use cleaning agents that contain a sives



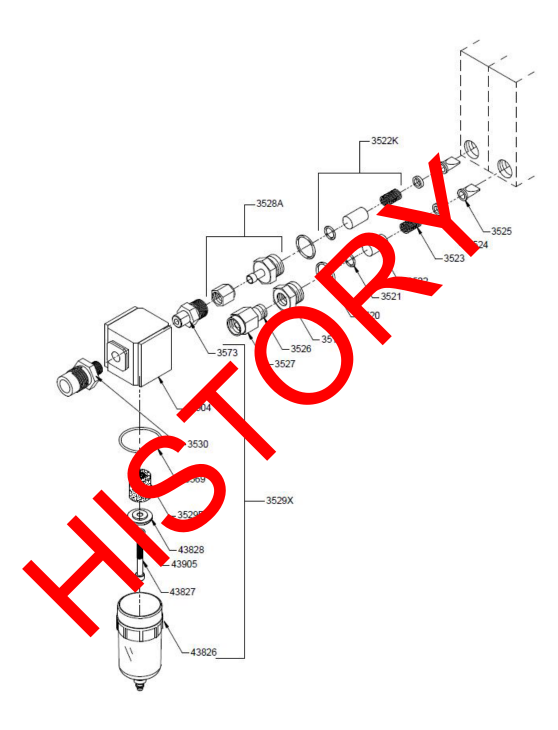


Figure 4

Factory overhaul

In order to assure proper function and accuracy, the Sechrist air/oxygen mixers must be thoroughly overhauled every two- (2) years. To maintain the product warranty, this overhaul must be performed by Sechrist Industries or by Sechrist authorized personnel.

TROUBLESHOOTING

Problem	Possible Cause	Sorrective Action
Inaccurate FIO ₂	O ₂ analyzer at of	Recalibrate O_2 analyzer.
	calibration (a pst co problem).	
	problem.	
	Inproper put of supply	Check/verify supply gas
	ses.	purity.
	Inc act or supplied to	Assure that outlets and
	inlet.	hoses are connected
		correctly.
	Front and rear seats are	**Clean or replace seats.
	worn.	
	Incorrect calibration of	**Recalibrate mixer as
	proportioning module.	outlined in the service
		manual.
	Malfunctioning balancing	**Recalibrate the balancing
	module.	module as outlined in the
		service manual.
FIO ₂ control knob is	Faceplate has shifted.	Reposition faceplate.
difficult to turn.		
	Bent adjustment shaft.	** Replace shaft and recalibrate as outlined in
		the service manual.
	Air or O_2 inlet filter may be	Replace inlet filter.
FIO_2 change $> 1\%$ when	dirty causing a > 20 psi	1

Problem	Possible Cause	Corrective Action
testing.	(138 kPa) difference.	
	Regulator needle out of calibration.	**Recalibrate mixer as outlined in the service manual.
Continuous alarm with both inlet pressures equal.	Dirty inlet filter(s). Bypass check ball leal	* Clean check ball and
	Alarm module case calibration.	**Recalibrate as outlined in the service manual.
Alarm not sounding with the loss of pressure from one source gas.	Aron modul, out of calibra.	**Replace alarm reed. **Recalibrate as outlined in the service manual.
	Alarm poppets stuck.	**Clean, lubricate poppets and recalibrate as outlined in the service manual.

the Ablem or concern continues after taking the appropriate corrective action, consult an authorized Sechrist service representative or contact Sechrist Industries Technical Support.

^{**} To be performed only by authorized personnel.

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