

BONHAWA

Respiratory Humidifier

Instructions for Use (IFU)

U.S. Version

Model: RHF G303

Caution: **Rx Only** Federal law restricts this device to sale by, or on the order of, a health care professional.

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Note: This product is protected by the following granted, continued, and pending patents: US 11,135,390; US 11,318,268; US 11,468,988; US 11,954,331; US 11,642,477; US 11,702,602, US 11, 826,510; US 11,865,259; US 11,896,847; etc.

1. Preface

The Instructions for Use (IFU) includes the information regarding set-up, operation, and

maintenance for the Bonhawa Respiratory Humidifier, model RHF G303. The QuickStart

Guide included in your packaging should be attached to its stand as a reference for

professional users. The Bonhawa Respiratory Humidifier is for non-invasive use (NIV) only.

This device shall be used by a healthcare professional. Read through these Instructions for

Use and QuickStart Guide before using the system.

Indication for use 2.

The Bonhawa Respiratory Humidifier is intended to provide highflow warmed and

humidified respiratory gases for administration to spontaneously breathing patients 10 kg

and up, child to adults in hospitals. It adds heat and moisture to the flow of gases. The flow

may be from 2 to 70 L/min depending on the patient interface.

The Bonhawa Respiratory Humidifier provides high flow gases with simultaneous oxygen

delivery to spontaneously breathing patients without bypassed upper airways in hospitals.

Contraindications 3.

The Bonhawa Respiratory Humidifier is contraindicated for use with unresolved tension

pneumothorax and facial trauma.

4. Product Name and Model

Product Name: Bonhawa Respiratory Humidifier

Product Models: RHF G303

RHF G3 03

- Third Identification Code for Product

Secondary Identification Code for Product

Product Platform Code for Bonhawa Humidifier

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5. Symbols

The following symbols may appear on the product or packaging.

	Read instructions before use	[i	Refer to instructions	
\triangle	Warning or caution		Warning: Hot Surface	
	Maximum water level	IP22	Ingress protection rating	
~~	Date of manufacture		Manufacturer	
LOT	Lot Number	SN	Serial number	
	Electronics waste to be disposed of properly	RoHS	Compliant with RoHS guidelines	
†	Type BF applied part	EC REP	EU Representative	
	Alarm Symbol		Alarm Paused	
	Audio Paused	$\overline{}$	Power On/Off	
	Class II equipment	REF	Model/ Catalog Number	
G	Settings Locked	4	Settings Unlocked	
\$	System Settings	NON	Non-sterile	
2	Do not re-use	R only	Federal law restricts this device to sale by or on the order of, a medical professional	
MD	Medical Device		MRI Unsafe	

6. Contents List

Name	Quantity	Name	Quantity
Bonhawa Respiratory Humidifier	1	Power Cord	1
Quick Start Guide	1	Air Filter	1

7. Warnings and Cautions

Warnings: Warning identify conditions or practices that could result in serious adverse reactions or potential safety hazards.

Cautions: Cautions identify conditions or practices that could result in damage to the humidifier or other equipment.

Warnings:

- The humidifier is not a life-support device.
- The humidifier uses O₂ concentration, respiratory gas flow and temperature, and provides alarms which need to be controlled and monitored to achieve its intended use.
 The humidifier shall be used only by healthcare professionals.
- Closely monitor the patient's heart rate, respiratory rate, and SpO₂ through pulse oximetry.
- The water chamber, water chamber adaptor, nasal cannula and breathing circuit are single patient use only.
- To prevent patient, bystander, and equipment harm, do not operate or place the humidifier in an MRI (Magnetic Resonance Imaging) environment.
- The performance of the humidifier may change if used in combination with devices other than the ones recommended by the manufacturer (e.g., nasal cannula).
- The humidifier is not intended to treat patients whose upper airways have been bypassed.
- Keep the power cord away from any surface with high temperature.

To avoid strangulation, arrange the power cord and the breathing circuit such that they

will not circle around the neck or head.

Only the accessories approved or recommended by Telesair should be used with the

humidifier. Safety and effectiveness may be compromised if using accessories that are

not approved or recommended by Telesair.

Stop using the humidifier and contact Telesair Customer Service if any of the following

occurs: unexpected noise, damaged device housing or any change in performance for

unknown reasons.

Do not immerse the device or accessories into fluid, including power cable or power

plug.

• Power off and unplug the humidifier prior to cleaning.

• Power off and unplug the humidifier immediately when liquids are spilled onto the

device. Let the device dry prior to plugging power back to the humidifier.

The surrounding area of the system shall be kept dry and clean.

Covering breathing circuits with a blanket or heating them in an incubator or with an

overhead heater can affect the quality of the therapy or injure the patient.

• Do not directly touch the heater plate by hand or body within 10 minutes after the

treatment.

• Avoid using the humidifier adjacent to or stacked with other equipment.

• If there is any flame in the vicinity, do not use the humidifier.

• The humidifier shall not be used in the operating room or in the presence of flammable

gases.

• Portable RF communications equipment (including peripherals such as antenna cables

and external antennas) should be used no closer than 30 cm (12 inches) to any part of

the Bonhawa humidifier System, including cables specified by the manufacturer.

Otherwise, degradation of the performance of this equipment could result.

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System performance may be adversely affected, and device may be damaged if incorrect

substances or methods for cleaning are performed.

The humidifier shall not be used with nitric oxide. Such use might cause the humidifier

to not function correctly causing serious deterioration of health.

When the humidifier is in operation, do not block the air intake port on the device or

occlude the breathing circuit.

• Place the device on a level surface, lower than the patient's head to prevent water from

entering the breathing circuit or patient interface when the humidifier is in operation.

Monitor the breathing circuit for condensate at low flow and lower temperatures. D

not drain condensate back into the humidifier chamber.

Choose an appropriate cannula size that does not occlude the nares. Occlusion of the

nares by the applied cannula can result in dermal damage and pulmonary barotrauma.

CAUTIONS:

Normal operation of the humidifier shall be verified if using the humidifier adjacent to

or stacked with other equipment is necessary.

The humidifier shall be used in an environment with good ventilation.

The Bonhawa humidifier should not be used adjacent to or stacked with other

equipment. If adjacent or stacked use is necessary, the Bonhawa humidifier should be

observed to verify normal operation. If operation is not normal, the Bonhawa humidifier

or the other equipment should be moved.

Use of accessories, transducers, and cables other than those specified or provided by

the manufacturer of this equipment could result in increased electromagnetic emissions

or decreased electromagnetic immunity of this equipment and result in improper

operation.

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LB0036 Revision E Page 8 of 39 Avoid exposure to known sources of EMI (electromagnetic interference) such as

diathermy, lithotripsy, electrocautery, RFID (Radio Frequency Identification), and

electromagnetic security systems such as anti-theft/electronic article surveillance

systems, metal detectors. Note that the presence of RFID devices may not be obvious. If

such interference is suspected, reposition the equipment, if possible, to maximize

distances.

To prevent water inside the water chamber from getting into the humidifier, do not place

the device upside down or on its side.

Ensure that the water chamber is empty or removed prior to transporting the humidifier

system.

• Allow the water chamber to cool down for at least 10 minutes before removing it from

the humidifier.

• System performance may be adversely affected, and device may be damaged if incorrect

substances or methods for cleaning are performed.

• Do not remove the humidifier cover. Any repair or servicing activity must be carried out

by Telesair-authorized service personnel using appropriate tools.

8. Disclosures

The Bonhawa Respiratory Humidifier is not made with natural rubber latex.

The Bonhawa Respiratory Humidifier is not made using phthalates including DEHP.

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9. Device Overview



Figure 9-1 Front and Back of Bonhawa Respiratory Humidifier

Table 9-1 General assembly components

1	Top Outer Housing	13	13 Water Chamber Adaptor Interface	
2	Alarm Light Indicator	14	Water Chamber	
3	Touch Screen Display	15	Water Chamber Adaptor	
4	Power Indicator	16	Heated Breathing Circuit	
5	Power On/Off Button	17	Power Cord	
6	Alarm Silence Button	18	Speaker	
7	Dial	19	Gas Inlet Cover (Oxygen supply & Air)	
8	Display Panel Cover	20	Oxygen Inlet Port	
9	Bottom Outer Housing	21	Thumb Screw	
10	Finger Guard	22	Label	
11	Heater Plate	23	Power Cord Retaining Clip	
12	Air Outlet			

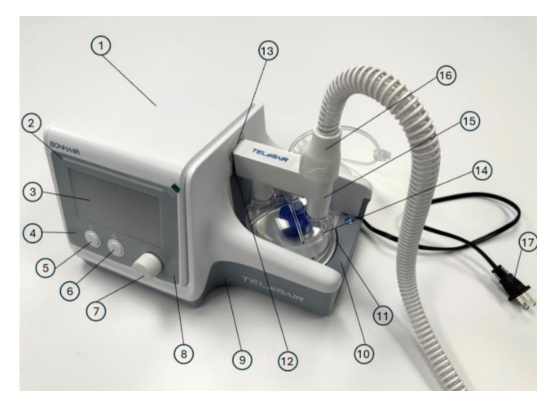


Figure 9-2 Feature Diagram of Bonhawa Respiratory Humidifier, Front View.



Figure 9-3 Feature Diagram of Bonhawa Respiratory Humidifier, Rear View



Figure 9-4 User Interface of Bonhawa Respiratory Humidifier

Table 9-2 Heated Breathing Set (HBCK01) Contents



Table 9-3 Patient interface consumable accessories

	Size Description	Catalog Number	Recommended Flow Rate Range
	C-mall	ENCO2	2 – 25 L/min (Pediatric Mode)
	Small	ENC03	10 – 40 L/min (Adult Mode)
	Medium	ENC02	10 – 70 L/min (Adult Mode)
Patient Interface, Nasal			
Cannula	Large	ENC01	10 – 70 L/min (Adult Mode)

Figure 9-5. Bonhawa optional Trolley (CTP1)



Table 9-4 Customer provided accessories not included with Bonhawa Humidifier but necessary for use.



DISS Fitting for Oxygen Flow Meter	Oxygen Flow Meter	O₂ Tubing (2m)	Sterile Distilled Water
------------------------------------	-------------------	----------------	-------------------------

The Oxygen Flow Meter provides a diameter indexed safety system (DISS) fitting that connects the hospital facility oxygen supply to the inlet of the Oxygen Flow Meter. The outlet of the Oxygen Flow Meter connects to a DISS fitting that provides a barbed connector interface that facilitates the Oxygen (O₂) Tubing connection to the O₂ inlet of the Bonhawa Respiratory Humidifier. Both the flowmeter and nipple are representative of the types of respiratory delivery components that are commonly used in hospital respiratory departments and long-term care facilities. **Note:** The Oxygen Flow Meter must have a flush flow rate of 80 LPM or higher to support the requirement of the humidifier.

The Oxygen Tubing is also a commonly used component available in hospital respiratory departments. Sterile distilled water is used to provide a humidification water source for the Bonhawa device. Sterile distilled water is used and is readily available in hospitals and long-term care facilities. This sterile water is provided in a bag or bottle that is to be spiked to provide a water line from the bag to the inlet of the water chamber. The water chamber is provided with a spike tubing connection for completing this fluid interface.

Refer to your hospital policy or contact your local sales representative for assistance in locating these accessories if they are not readily available in your facility.

10. Assembly

Getting Started

- ✓ Identify the necessary components.
- ✓ Place the humidifier on a level and secure surface, which is slightly lower than the head of the patient.
- Open the bag containing the heated breathing set (breathing circuit, water chamber and water chamber adaptor).

2. Water Chamber Insertion:

✓ Remove the caps on the water chamber, connect the water chamber adaptor to

the water chamber, push the assembly all the way down onto the ports of the

water chamber. Make sure that there is a tight connection between the water

chamber adaptor and the water chamber.

✓ To install the water chamber onto the humidifier, firmly press down the finger

guard and slide the water chamber towards the humidifier until the inlet port of

the water chamber adaptor is firmly connected with the humidifier and the finger

guard is up and in locked position.

3. Water Source Installation

✓ Place the water tubing into the clip present on the water chamber adaptor.

✓ Hang a Sterile bag of distilled water about 20 cm (8") above the humidifier,

connect the water chamber tubing via spike port into the water bag.

✓ Open the cap on the spike port, the water from the bag should enter the drip

chamber then fill the water chamber automatically.

Check the water chamber and make sure it does not exceed the max water level.

MAXIMUM WATER LEVEL

NOTE: Make sure that both the water bag and the water chamber are not empty

during operation to ensure continued humidification.

WARNING: Adding substances other than sterile distilled water can adversely affect the

normal functionality of the humidifier

4. Breathing Circuit Installation

✓ To install the heated breathing circuit, push all the way down to connect one end

of the breathing circuit (with the tear drop shape connector) to the vertical outlet

port of the water chamber adaptor, double check if the heated breathing circuit is

securely fastened onto the water chamber adaptor.

5. Patient Interface Installation

Connect the patient interface, such as a nasal cannula, to the other end of the

breathing circuit.

11. Connect to Oxygen Supply

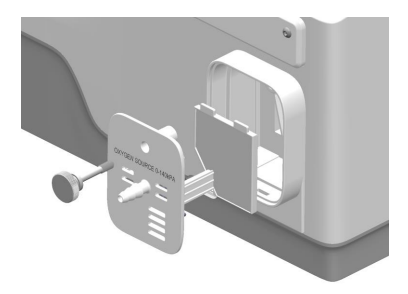


Figure 11-1 Air Filter Assembly Overview

- 1. Check if the Air Filter is installed properly.
- 2. Secure the O₂ Inlet to the device with the thumbscrew.
- 3. Power ON the device.
- 4. Connect the O_2 tubing to the O_2 inlet port and ensure that the connection is tight and secure.
- 5. Verify the $O_2\%$ displays 21% on the display when there is no oxygen flowing through the device.

WARNING: Do not connect the Oxygen supply before turning on the device.

- 6. Turn ON oxygen supply and start therapy based on the descriptions in Section 12.2
- 7. Observe the $O_2\%$ display increase to the intended Oxygen concentration level.
- 8. Turn OFF oxygen supply when therapy is completed.
- 9. Power OFF the device.

12. Operation

12.1. Start Therapy

1. The operator must check if the entire humidifier system has been correctly set up

for the patient who will be receiving the treatment, including therapy setting

parameters and the accessories used. To help achieve better efficacy, the operator

should assess the setting parameters and alarm limits periodically during the therapy.

WARNING: Operator shall make sure that all parts or "accessories" used are manufacturer

approved and are compatible with the humidifier.

WARNING: Using accessories that are not approved by Telesair may compromise the safety

and efficacy of the device. Use only the breathing circuits and accessories that are approved

by Telesair with the Bonhawa humidifier.

NOTE: The accessories recommended or approved by Telesair can be found in Table 9-

2 and Table 9-3.

WARNING: Before use, the operator must make sure that the power cord is in good

condition, and a good power connection between the device and the power source is

secured during operation.

2. Connect the humidifier to main power using the power cord provided. Ensure the

latch is engaged to prevent the power cord disconnecting inadvertently.

3. To power on the humidifier, press the Power On/Off Button on the humidifier control

panel. The therapy page will be displayed within 15s when powered on and the

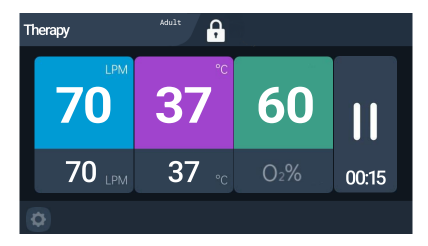
system enters Standby.

4. Do not initiate the flow of oxygen into the device until the Bonhawa Respiratory

Humidifier is in Therapy Mode.

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5. Press the dial once, *Therapy* starts. On the screen, the flow, temperature, and oxygen concentration are displayed.



6. Initiate oxygen flow using the external flow meter to achieve the desired O₂ concentration.

12.2. Oxygen Concentration

The $O_2\%$ displayed on the device screen is the Oxygen concentration level during therapy delivered to the patient. Regularly check the displayed $O_2\%$ and the desired SpO_2 level while the Bonhawa Respiratory Humidifier is delivering therapy.

12.3. Stop Therapy

- 1. Turn off the flow of oxygen before switching from Therapy Mode.
- 2. To stop the therapy, press down the Dial key during *Therapy*, the humidifier will stop and begin to cool down.
- 3. When system is cooled down, the humidifier will enter Standby.
- 4. In Standby, press and hold the Power On/Off Button for 4-5 seconds to initiate power off cycle, power off the humidifier by pressing Confirm on the touch screen.

13. Setting Functions

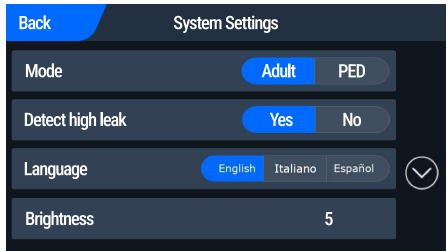
The settings of the humidifier should only be adjusted by trained healthcare professionals.

13.1. Enter the Setting Functions

1. Press Lock Key icon on the touch screen to unlock it.

- 2. Once the screen is unlocked, therapy settings can be changed.
- 3. To change the therapy mode (Adult or Pediatric), press the *Gear* icon to enter *System Settings* page.





4. Once therapy setting changes are completed, press the *Lock* icon to save and apply the settings.

13.2. Set Flow

- On the setting page, the flow setting can be entered by pressing the Flow (LPM) on the touch screen. The upper part of the icon will turn grey, and the lower values of the icon can be adjusted.
- 2. Rotate the Dial key to adjust the setting value to adjust the flow setting.
- 3. Push the Dial key or touch the Flow setting to confirm the setting value.

13.3. Set Gas Delivery Temperature

- At the setting page, the gas delivery temperature parameter can be entered by pressing Temperature.
- 2. Rotate the dial to adjust the setting value to adjust the gas delivery temperature setting.
 - Adult mode: the gas delivery temperature can be set to 31°C, 34°C, or 37°C
 - <u>Pediatric mode:</u> the gas delivery temperature can be set to 34°C
- 3. Push the Dial to confirm the setting value.

13.4. Set Threshold for Oxygen Concentration Alarm

The thresholds for O_2 too High and O_2 too Low alarms can be adjusted on the *Oxygen Alarm Limit Setting* page.

- 1. On the setting page, press the $O_2\%$ on the display to enter the *Oxygen Alarm Limit Settings* page.
- 2. Press the Oxygen Upper Limit setting, the setting value is highlighted.
- 3. Use the Dial to change the limits.
- 4. Press the limit value to accept the change.
- 5. Press Back to exit the Oxygen Alarm Limit Setting page.
- 6. Press the *Lock* icon to save the settings.

WARNING: Setting O_2 alarm limits at extreme thresholds will effectively disable O_2 alarm detection.

13.5. Set Therapy Mode

Based on the patient who will be receiving the treatment, the therapy mode can be set to Adult Mode or Pediatric Mode. The therapy can be set on the *System Settings* page.

- 1. Press the Lock Key icon.
- 2. Press the Gear icon to enter System Settings page.
- 3. Select Adult or PED therapy mode.

4. Press Back.

5. Press the *Unlock Key* icon to save the setting.

13.6. Set High Leak Detection

When using a patient interface with low resistance other than the supported cannulas, "Check Leaks" alarm may persist. To stop the undesired alarm, high leak detection can

be turned OFF.

1. Press the Lock Key icon.

2. Press the *Gear* icon to enter *System Settings* page.

3. Select Yes or No to detect or not detect high leak.

4. Press Back.

5. Press the *Unlock Key* icon to save the setting.

WARNING: Turning off high leak detection will not alarm when the cannula is disconnected. Only turn off high leak detection when needed.

13.7. Set Brightness

The screen brightness can be set on the System Settings page.

1. Press the Lock Key icon.

2. Press the Gear icon to enter System Settings page.

3. Press the *Brightness Value*, the setting value is highlighted.

4. Use the dial to change to Brightness Value.

5. Press the Brightness Value to save the setting.

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13.8. Factory Default Settings

Therapy mode Adult

Flowrate 30LPM

Temperature 34°C

O₂ alarm lower limit 21%

O₂ alarm upper limit 95%

Detect high leak Yes

Brightness 8

13.9. Settings Storage

All settings are saved and are maintained across power cycles.

14. Alarms

The device generates an alarm with both visual and audio indications to alert the user that the normal operation of the device is interrupted by conditions listed in Table 13-1. The table lists Alarm priority, Alarm Message, its description (with fault detection), impact and fault responses. The alarm priority is listed in descending priority; lower priority number is higher in alarm priority, with one being the highest priority. System error alarms are technical alarms. All other alarms are patient alarms.

Depending on the alarm condition, alarm detection may take up to 120 seconds to trigger an alarm.

A visual alarm message will be displayed on the screen and the corresponding audio alarm is a repeating 3-beep sound for every 5 seconds. By pressing down the audio pause key, the audio alarm signal can be silenced for 2 minutes.

Alarm settings are retained and restored after shutdown or power loss.

Table 14-1 List of Alarm Conditions

Alarm Priority	Alarm Messages	Conditions	Impacts	Response Actions
				Turn off the
	"System Error		Normal	device and try to
	nnn"	Internal fault is	device	trouble shoot as
1		detected by the		per the technical
	"Please Turn Off the	device	operation is	manual; unplug
	device"		interrupted.	the device if
				necessary
		The water		Connect the
		chamber		Water Chamber
2	"Check Water Chamber	adaptor is not	Flow and O ₂	Adaptor properly
2	Adaptor"	installed	1 low and 02	or replace if alarm
		correctly or is		persists
		faulty		pc131313
		The breathing		Connect the
		circuit is not		Breathing circuit
2	"Check Circuit"	installed	Flow and O ₂	properly or
		correctly or is		replace if alarm
		faulty		persists
				Check for leaks in
		The system has		the device/
2	"Check Leaks"	high flow	Flow and O ₂	patient interface
		leakage		connection and
				resolve if needed.
				Check for the
		The circuits or		occlusion in the
2	2 "Circuit Occlusion"	patient's user	Flow and O ₂	circuit and
		interface is	1 low and O2	interface and
		occluded		remove the
				occlusion

Alarm Priority	Alarm Messages	Conditions	Impacts	Response Actions
				Check the
		The oxygen level		external O ₂
2	"Oxygen too Low"	is below the	O ₂ Levels	flowrate settings
		threshold		and the O ₂ alarm
				settings
				Check the
		The oxygen level		external O ₂
2	"Oxygen too High"	is above the	O ₂ Levels	flowrate settings
		threshold		and the O ₂ alarm
				settings
2	"Flow Too Low"	The flow cannot reach the flow setting	Flow, O ₂ and humidity levels	Restart the device
2	"Flow Too High"	The flow is more than the set value	Flow, O ₂ and humidity levels	Restart the device
2	"Check Water Chamber"	The water ran out in the water chamber	Humidity level	Stop the therapy, replace the water bag and restart the therapy.
2	"Temperature Too Low"	The gas temperature cannot reach the setting value	Humidity level	Restart the device.
2	"Temperature Too High"	The gas temperature exceeds the setting value	Humidity level	Restart the device.

Alarm Priority	Alarm Messages	Conditions	Impacts	Response Actions
			Normal	
0	No message – Audio alarm only	The power is disconnected while the device is ON	device operation is interrupted. Device is powered off.	Check power and the power connection

14.1. Alarm Signal Functionality Testing Method

The Alarm Signal Functionality can be tested by following the steps given below.

NOTE: This test can be performed at any time when the device is turned on and off patient.

- 1. Once the device is installed properly and ready to be used, Turn on the device.
- 2. Remove the breathing circuit.
- 3. Verify the "Check Circuit" visual alarm signal, displayed alarm message, and alarm sound is audible.

CAUTION: Do not use the humidifier if either of the alarm indications is absent. Refer to the Bonhawa Respiratory Humidifier technical manual for troubleshooting procedure. If the problem persists, please contact Telesair Customer Service associate.

15. Maintenance

It is important that humidifier is cleaned between patients as well as on a weekly basis during normal use and/or for same patients to ensure best treatment. Follow the instructions in the sections below to learn how to disassemble, clean, inspect and reassemble your device.

The manufacturer will provide the schematics, parts list, and other documents to facilitate maintenance. Please contact Telesair Customer Service for questions about maintenance.

15.1. Inspection and Replacement Schedule

WARNING: All the consumables are single patient use only. They must be disposed of after use and between patients.

Parts/ Accessories/ Consumables	Inspection	Maximum use/Replace after
		4 weeks
		WARNING: Replace the
Air Filter	Every week	filter immediately if it is
		damaged or granular dust
		clogging is observed
Heated Breathing Set		
(Water Chamber Adaptor)		1 Week/every patient
(Breathing Circuit)	Every Use	
(Water Chamber)		
Nasal Cannula		1 Week/every patient

15.2. Disassembling and Replacement Procedure

Air Filter

- ✓ Power OFF the device.
- ✓ Disconnect oxygen tubing.
- ✓ Unplug the Power Cord.
- ✓ Unscrew the Thumb Screw.
- ✓ Remove the O₂ Inlet Cover.
- ✓ Remove the old Air Filter and install a new Air Filter and close the cover.

Water Chamber Adaptor/ Breathing Circuit/ Water Chamber/ Nasal Cannula

- ✓ Turn OFF the device.
- ✓ Disconnect the Power Cord.
- ✓ Firmly hold Water Chamber Adaptor with one hand and pull the breathing circuit in upward direction with the other hand.
- ✓ Carefully press down the Finger Guard and pull out the Water Chamber along with the Water Chamber Adaptor.
- ✓ Replace with new accessories as needed.

15.3. Exterior and Touch Panel Cleaning

- **CAUTION:** To prevent possible damage to the humidifier, use only those cleaning and disinfecting agents listed in this document.
- CAUTION: To prevent possible damage to the humidifier, do not drip or spray
 any liquids directly onto any surface including the front touch panel,
 keys, knobs, connection points, and ports.
- CAUTION: Never clean or disinfect the touch panel with an abrasive brush or device, as this will cause irreparable damage.
- **CAUTION:** Do not attempt to sterilize or autoclave the humidifier.
- **CAUTION:** Do not attempt to submerge or wash humidifier with excessive liquids.
- **CAUTION:** Do not attempt to clean or disinfect humidifier while the heater plate is hot.
- CAUTION: Allow the humidifier to dry prior to reconnecting the Power Cord
- **NOTE:** Use of unapproved cleaning and disinfecting agents may cause

damage to the enclosure, touchscreen, or parts of the humidifier.

- NOTE: Do not clean or disinfect the internal area of the air outlet port, as this
 can damage the valve located within this port.
- NOTE: Standard cleaning and disinfection process must be performed on the humidifier between patients.

The water chamber, water chamber adaptor, breathing circuit, and the patient interface are consumables, which must be changed between patients or changed every week for the same patient.

15.3.1. Approved Cleaning Agents

The following cleaning agents are acceptable for use on the front touch panel and exterior surfaces of the humidifier.

Cleaning Agent Ruhof Endozime® Multi-Tiered Enzymatic Detergent or equivalent

15.3.2. Cleaning Instructions

- 1. Power off and unplug the humidifier before cleaning.
- 2. The operator must wash his/her hands properly and avoid touching the connection port without gloves.
- 3. Apply cleaning agent to a soft lint-free cloth or use a disposable wipe. The cloth or wipe should be saturated, not dripping.
- 4. Wipe cleaning agent over the entire exterior surface and touch panel of humidifier.
- 5. Let the cleaning agent remain on the surface for 90 seconds.
- 6. Continue wiping until all visible contaminates and soiling are removed.
- 7. Rinse with a clean, water-dampened cloth for 90 seconds and let dry completely before reusing.

15.4. Exterior and Touch Panel Disinfecting

- **CAUTION:** To prevent possible damage to the humidifier, use only those cleaning and disinfecting agents listed in this document.
- CAUTION: To prevent possible damage to the humidifier, do not drip or spray
 any liquids directly onto any surface including the front touch panel, keys,
 knobs, connection points, and ports.

- CAUTION: Never clean or disinfect the touch panel with an abrasive brush or device, as this will cause irreparable damage.
- **CAUTION:** Do not attempt to sterilize or autoclave the humidifier.
- **CAUTION:** Do not attempt to submerge or wash the humidifier with excessive liquids.
- **CAUTION:** Do not attempt to clean or disinfect the humidifier while the heater plate is hot.
- **CAUTION:** Allow the humidifier to dry prior to reconnecting the Power Cord
- **NOTE:** Use of unapproved cleaning and disinfecting agents may cause damage to the enclosure, touchscreen, or parts of the humidifier.
- NOTE: Do not clean or disinfect the internal area of the air outlet port, as this
 can damage the valve located within this port.
- NOTE: Standard cleaning and disinfection process must be performed on the humidifier between patients.

The water chamber, water chamber adaptor, breathing circuit, and the patient interface are consumables, which must be changed between patients or changed every week for the same patient.

15.4.1. Approved Disinfecting Agents

The following disinfecting agents are acceptable for use on the front touch panel and exterior surfaces of the humidifier.

Disinfectant
CLOROX Disinfecting Wipes or equivalent

15.4.2. Disinfecting Instructions

- 1. Power off and unplug the humidifier before cleaning.
- 2. The operator must wash his/her hands properly and avoid touching the connection port without gloves.
- 3. Utilize disinfecting agent wipes and apply over the entire exterior surface of the humidifier for 90 seconds.
- 4. Let the disinfectant remain on the surface for the contact times indicated in the specifications for the disinfecting agent.
- 5. Rinse with a clean, water-dampened cloth for 90 seconds and let dry completely before reusing.

15.5. Service

The Bonhawa Respiratory Humidifier is designed to operate without scheduled preventative maintenance service. Under normal conditions, periodic service is not required. If the humidifier has been subjected to abnormal conditions (e.g., high impact such as a hard drop), service is needed to verify performance and safety. This shall include a gas pathway leak test. Contact your local Telesair representative with questions regarding service and operation of your Bonhawa Respiratory Humidifier or email service@telesair.com.

Waste Disposal

16.1. Accessories

When expended, the disposable water chamber, water chamber adaptor, breathing circuit and patient interface must be disposed based on local regulations for waste management.

16.2. The Device

Consult with the local waste management services or your distributor before recycling the device. Special handling and disposal for electrical or electronic equipment waste may be required based on local regulations.

17. Troubleshooting

Read the following table for troubleshooting when the humidifier system is not working as intended. Contact Telesair Customer Service if you are not able to solve the problem. Do not open the enclosure of the humidifier at any point.

Table 17-1 Troubleshooting

Problem	Possible Cause	Trouble Shooting Actions
	Power to the humidifier	Connect the humidifier to power source,
Screen is off	might have been	making sure there is a good and secure
	disconnected	connection between the two.

Condensation of water	The ambient	
vapor inside the	temperature might be	Try to increase the ambient temperature
patient's nose, the	too low	
breathing circuit, or the	The setting of humidity	Change the humidity setting to a lower
patient interface	level might be too high	level

18. Technical Specifications

18.1 Input Specification

Model	RHF G303
Power Supply AC Voltage	100-120V
Power Supply Frequency	50/60Hz
Power Supply Current	1.2A (2.4A max)
Maximum Oxygen Gas Supply	70 L/min max

18.2 Air Filter

Material:	Double Laminated Spunbond Polypropylene
	>99.5% at 32LPM (0.1 micron)
Efficiency:	99.9996% Bacterial Filtration (40cm²)
	99.996% Viral Filtration (40cm²)

18.3 Operating Environment

Temperature	+18°C to +28°C
	10 % to 85 %, non-condensing, but not
Humidity	requiring a water vapor partial pressure
	greater than 50 hPa
Altitude	0 - 2000 m

WARNING: Do not use the humidifier at altitude higher than 2000 m (6000 ft) or outside the temperature range of +18°C to +28°C. The quality and efficacy of therapy can be adversely affected.

Environmental Conditions for transport or Storage: –10°C to +60°C, at a relative humidity between 10% to 95%, non-condensing.

Classification			
Type of protection against electric shock	Class II Equipment		
Degree of protection against electric shock	Type BF Applied Part		
Degree of protection against ingress of water	IP22		
The degree of safety when used under			
flammable anesthetic gas mixed with air			
or flammable anesthetic gas mixed with	Non-AP/APG type		
oxygen or nitrous oxide			
Operating mode	Continuous		
Phy	rsical		
Dimension (H x W x D)	319 mm X 223 mm X 185 mm		
Weight	2.6 Kg		
Water capacity	450 + 00 - 1		
(When filled to maximum water level)	150 ± 30 mL		
Pressure drops across water chamber	<1 hPa, when flow rate is 60 LPM,		
Water chamber leak	<0.025 LPM, when pressure is 60hPa		
	<0.7 mL/hPa when empty		
Water chamber compliance	<0.5 mL/hPa, when at maximum water		
	level		
Maximum working pressure of humidifier	40hPa		
Sound Pressure Level does not exceed	50 dB(A)		
during normal operation	30 db(A)		
Temperature			
Warm-up time needed, when starting	10 min to reach 31°C and 34°C		
temperature is 23±2°C:	30 min to reach 37°C		
Maximum temperature of delivered gas	≤43°C		
Oxygen Monitor			
Accuracy	≤±3%, between 21% and 95%		
Humidification Performance			
37°C	≥33mg/L		

Classification		
34°C	≥16mg/L	
31°C	≥16mg/L	

NOTE: The displayed temperature is referenced at the outlet of the water chamber adaptor.

Therapy Mode	Flow Setting Range	Flow Setting Resolution	Flow Delivery Accuracy
		1 LPM for flow range	
Adult	10 to 70	between 10 and 25 LPM.	
Mode	LPM	5 LPM for flow range	+150/ of roading or 2 LDM
		between 25 and 70 LPM	±15% of reading or 3 LPM,
Pediatric Mode	2 to 25 LPM	1 LPM	whichever is greater

NOTE:

- Flow rate is expressed in BTPS condition in this document.
- The performance of humidification may decrease by the presence of large unintended leaks.

Alarm

The alarm system complies with IEC 60601-1-8, Medical electrical equipment – Part 1-8: General requirements for basic safety and essential requirements; collateral standard - alarm systems.

Alarm sound level exceeds 45 dB(A) @ 1m.

Cables that are likely to affect the EMC compliance are listed in the following table.

Туре	Maximum length	
Power cord	2.0 m	

NOTE: Use only Telesair-provided power cords

Electromagnetic compatibility (EMC)

The BONHAWA HUMIDIFIER is suitable for the electromagnetic environment of typical hospital or long-term care facilities.

During the immunity testing described below, the Bonhawa humidifier will continue to humidify respiratory gases within specification.

Manufacturer's declaration - electromagnetic emissions

The Bonhawa Respiratory Humidifier is intended for use in the electromagnetic environment specified below. The customer or the user of the Bonhawa Respiratory Humidifier should assure that it is used in such an environment

Emissions test	Compliance	Electromagnetic Environment - Guidance
RF emissions CISPR 11	Group 1	The BONHAWA HUMIDIFIER 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.
RF emissions CISPR 11	Class A	The BONHAWA HUMIDIFIER is suitable for use in all establishments other
Harmonic emissions IEC 61000-3-2	N/A	than domestic and may be used in domestic establishments and those directly connected to the public low-voltage power supply.
Voltage fluctuations / flicker emissions IEC 61000-3-3	N/A	network that supplies buildings used for domestic purposes, provided the following warning is heeded: Warning: This equipment/system is intended for use by healthcare professionals only. This equipment/ system may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating the BONHAWA HUMIDIFIER or shielding the location.

MANUFACTURER'S declaration – electromagnetic IMMUNITY – for all ME EQUIPMENT and **ME SYSTEMS**

Manufacturer's declaration - electromagnetic Immunity

The Bonhawa Respiratory Humidifier is intended for use in the electromagnetic environment specified below. The customer or the user of the Bonhawa Respiratory Humidifier should assure that it is used in such an environment

Immunity Test	Compliance level	Electromagnetic Environment - Guidance		
Electrostatic discharge (ESD) IEC 61000-4-2	±8 kV contact ±2, 4, 8 and 15 kV air	Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.		
Electrical fast transient/burst IEC 61000-4-4	±2 kV for power supply lines	Mains power quality should be that of a typical commercial or hospital environment.		
Surge IEC 61000-4-5	±1 kV differential mode	Mains power quality should be that of a typical commercial or hospital environment.		
Voltage dips, short interruptions, and voltage variations on power supply input lines IEC 61000-4-11	0 % 0.5 Periods 0% 1 Period 70% 25 Periods 0% 5 sec	Mains power quality should be that of a typical commercial or hospital environment. If the user of the BONHAWA HUMIDIFIER requires continued operation during power mains interruptions beyond that provided by the battery, it is recommended that the BONHAWA HUMIDIFIER is powered from an uninterruptible power supply.		
Power frequency (50/60Hz) magnetic field IEC 61000-4-8	30 A/m 50/60 Hz	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.		
Proximity magnetic fields IEC 61000-4-39	(8A/m, CW, 30kHz) (65 A/m (rms), PM at 2.1 kHz PM, 50% duty cycle, 134.2kHz) (75 A/m (rms), PM at 50 kHz, 50% duty cycle, 13.56MHz)	Avoid exposure to known sources of EMI (electromagnetic interference) such as diathermy, lithotripsy, electrocautery, RFID (Radio Frequency Identification), and electromagnetic security systems such as anti-theft/electronic article surveillance systems, metal detectors. Note that the presence of RFID devices may not be obvious. If such interference		

Manufacturer's declaration - electromagnetic Immunity				
The Bonhawa Respiratory Humidifier is intended for use in the electromagnetic environment specified				
below. The customer or the user of the Bonhawa Respiratory Humidifier should assure that it is used in				
such an environment				
	is suspected, reposition the equipment, if possible,			
		maximize distances		
NOTE U_T is the A.C. main voltage prior to application of the test level.				

MANUFACTURER'S declaration – electromagnetic IMMUNITY – for ME EQUIPMENT and ME SYSTEMS that are Professional Healthcare Facility Environment.

Manufacturer's declaration - electromagnetic Immunity

The Bonhawa Respiratory Humidifier is intended for use in the electromagnetic environment specified below. The customer or the user of the Bonhawa Respiratory Humidifier should assure that it is used in such an environment

Immunity Test	Compliance level	Electromagnetic Environment Guidance
Conducted RF	3 V 0.15 MHz – 80 MHz	
IEC61000-4-6		The BONHAWA HUMIDIFIER is suitable
	6 V rms in ISM bands	for the electromagnetic environment
Radiated RF		of typical hospital settings.
IEC61000-4-3	3 V/m 80 MHz to 2.7 GHz	

The BONHAWA Respiratory Humidifier was also tested for radiated immunity to RF wireless communication equipment at the test levels below.

Frequency (Hz)	Modulation	Level V/m
385	Pulse, 18 Hz, 50% DC	27
450	FM, 1 kHz Sine, ±5 Hz Deviation	28
710, 745, 780	Pulse, 217 Hz, 50% DC	9
810, 870, 930	Pulse, 18 Hz, 50% DC	28
1720, 1845, 1970		28
2450	Pulse, 217 Hz, 50% DC	28
5240, 5500, 5785		9

19. Warranty

The Bonhawa RHF G303 has a useful life of 5 years. Telesair will provide a warranty on the humidifier from the date of purchase for the time period listed below, if the device is used under normal conditions.

Model Type	Warranty Period
RHF G303	2 Years

Telesair will be responsible for repairing or replacing the defective product or any of its components if the product fails under normal conditions. This warranty is not transferable. It is only available to the original buyer of the device.

The coverage of the limited warranty will be lost if any of the following conditions occur:

- Repairs or services performed by any unauthorized service agency or personnel.
- Any damage resulted from abuse, modification, alteration, or any other improper use of the product.
- If the product is sold or resold outside the region of its original purchase.
- Any damage or contamination caused by cigarettes or other smoke.
- Warranty claims must be made by the original buyer of the device. Contact Telesair dealer or Telesair Customer Service for more information on warranty rights.

After Sales Service: Telesair Inc.

Registrant: Telesair Inc.

Address: 199 Technology Drive, Suite 110, Irvine, CA 92618, USA

Tel: +1-949-570-3553

Email: service@telesair.com

Web: https://www.telesair.com/

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Web: www.obelis.net