Tec 7 Vaporizer User's Reference Manual



User Responsibility

This Product will perform in conformity with the description thereof contained in this User's Reference manual and accompanying labels and/or inserts, when assembled, operated, maintained, and repaired in accordance with the instructions provided. This Product must be checked periodically. A defective Product should not be used. Parts that are broken, missing, plainly worn, distorted, or contaminated should be replaced immediately. Should repair or replacement become necessary, Datex-Ohmeda recommends that a telephonic or written request for service advice be made to the nearest Datex-Ohmeda Customer Service Center. This Product or any of its parts should not be repaired other than in accordance with written instructions provided by Datex-Ohmeda and by Datex-Ohmeda trained personnel. The Product must not be altered without the prior written approval of Datex-Ohmeda. The user of this Product shall have the sole responsibility for any malfunction which results from improper use, faulty maintenance, improper repair, damage, or alteration by anyone other than Datex-Ohmeda.

△ CAUTION

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U.S. Federal law restricts this device to sale by or on the order of a licensed medical practitioner. Outside the U.S.A, check local laws for any restriction that may apply.

Datex-Ohmeda products have unit serial numbers with coded logic which indicates a product group code. The year of manufacture and a sequential unit number for identification.

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	This alpha character indicates the year of product manufacture and when the serial number was assigned; "C" = 1999, "D" = 2000, "E" = 2001, etc. "I" and "O" are not used.

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Warranty

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1 Introduction

Precautions

⚠ WARNING

Do not fill the vaporizer with any agent other than the agent specified on the front label. The vaporizer is designed for that agent only. If any substance other than that specified is used, patient injury could occur.

United States (U.S.) Federal law restricts this device to sale by or on the order of a licensed medical practitioner. Outside the U.S., check local laws for any restrictions that may apply.

Do not attempt to use a vaporizer that has been dropped. A dropped vaporizer MUST be sent to the nearest Datex-Ohmeda Field Operations Unit for servicing.

Do not use malfunctioning equipment. Make all necessary repairs or have the equipment serviced by an authorized Datex-Ohmeda service center. After repair, test the equipment to ensure that it is functioning properly in accordance with the manufacturer's published specifications.

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Important

European Standard EN 740 - *Anesthetic Workstations and Their Modules* requires that an appropriate gas monitor is used to monitor the concentration of anesthetic agent vapor in the inspiratory gas when the vaporizer is in operation in order to provide protection against hazardous output in the event of a device malfunction.

Datex-Ohmeda strongly recommends the use of anesthesia gas monitoring with this equipment. Refer to local standards for mandatory monitoring.

Requests for servicing facilities, advice or assistance must be addressed to a local Datex-Ohmeda office.

Additional copies of this manual, can be requested from a local Datex-Ohmeda Field Operations Unit or a Datex-Ohmeda Authorized Distributor.

Datex-Ohmeda strongly recommends that you keep all relevant documentation, including this manual and accompanying labels, immediately available to all users.

Symbols

Warnings and Cautions tell you about conditions that can occur if you do not follow all instructions in this manual.

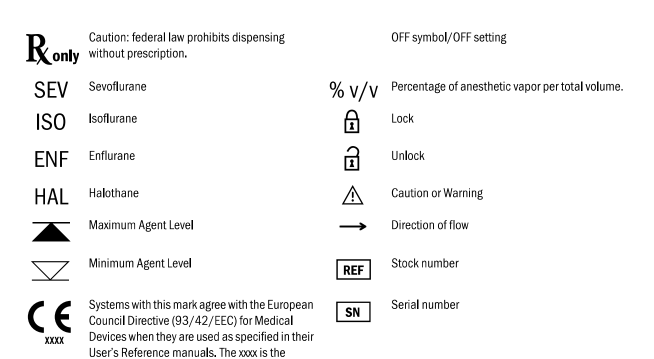
⚠ WARNING

Warnings tell about a condition that can cause injury to the operator or the patient.

⚠ CAUTION

certification number of the Notified Body used by Datex-Ohmeda's Quality Systems.

Cautions tell about a condition that can cause damage to the equipment. Read and follow all warnings and cautions.



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2 Description

What is a Tec 7 Vaporizer?

The Tec 7 Vaporizer is designed for use in continuous flow techniques of inhalation anesthesia. Each vaporizer is agent specific and is clearly labeled with the anesthetic agent that it is designed for.

The vaporizer is temperature, flow and pressure compensated so that its output remains relatively constant despite cooling due to evaporation, variations in inlet flow and fluctuating pressures as described in Section 7, Effects of Variables.

The vaporizer is designed to be used on Selectatec® Series Mounted Manifolds. The vaporizer can be installed on other Selectatec Manifolds but the interlock system is designed to function on Selectatec Series Mounted Manifolds only. Mounting a Tec 7 Vaporizer on a Selectatec 7 Compatibility Block is not recommended.

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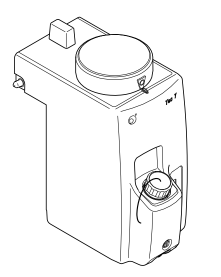


Figure 2-1 • Tec 7 Vaporizer

⚠ WARNING

Improper use may result in patient injury.

This manual and its associated documentation must be studied before any attempt is made to install, operate or clean any part of the Tec 7 Vaporizer.

The performance of the anesthesia machine and vaporizer can be degraded if the machine and vaporizer are mis-matched.

Only operate the vaporizer with dry medical gases.

If a vaporizer containing agent in the sump has been inverted, connect it to a gas scavenging system, set the dial to 5% and purge the vaporizer with the carrier gas at 5 liters/minute for 5 minutes.

⚠ CAUTION

The vaporizer is intended to be operated in its upright position.

Turn the vaporizer to $\ensuremath{\overline{\bigvee}}$ when it is not in use.

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Components

Control dial

A single control dial with a concentration scale calibrated in percentage of an esthetic agent vapor per total volume (% v/v) sets the desired concentration of the an esthetic agent.

A dial release in the dial assembly helps prevent accidental displacement of the control dial from the position. To select an ON setting, squeeze the dial release and simultaneously rotate the dial counter-clockwise.

The dial and dial release are designed to enable an ON setting to be selected using only one hand.

Safety interlocks

The vaporizer incorporates an interlock mechanism. This mechanism also interfaces with the Selectatec® Series Mounted Manifold to help satisfy the following criteria:

- The vaporizer must be locked onto the manifold before it can be turned ON.
- Only one vaporizer at a time can be turned ON when two or more vaporizers are fitted on a Selectatec® Series Mounted Manifold.
- The gas flow enters only the vaporizer that is turned ON.
- Any unwanted anesthetic trace vapor is minimized after a vaporizer is turned to
 .

⚠ WARNING

Earlier versions of the Selectatec Series Mounted Manifold that provide mounting positions for three vaporizers require that if only two vaporizers are fitted, then the center position must be occupied. If the center position is not occupied, the interlock that helps ensure that only one vaporizer at a time can be turned ON is ineffective.

Later versions of the Selectatec Series Mounted Manifold that provide mounting positions for three vaporizers incorporate an additional interlock that helps ensure that only one vaporizer at a time can be turned ON even if the center position is not occupied.

Vaporizer identification label

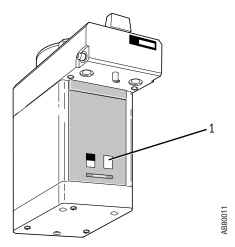
A vaporizer identification label is affixed to the back panel of the vaporizer as illustrated on Fig. 2-2.

An anesthesia system fitted with a vaporizer identification unit uses this label to identify the vaporizer type.

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⚠ WARNING

Do not affix any additional labels or markings to the back panel. They may adversely affect the operation of the vaporizer identification unit.



1. Vaporizer identification label

Figure 2-2 • Vaporizer identification label

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3 Setup and Mounting Procedure

Vaporizer mounting procedure

The vaporizer is designed to be used on Selectatec Series Mounted Manifolds. The vaporizer can be installed on other Selectatec Manifolds but the interlock system is designed to function on Selectatec Series Mounted Manifolds only.

Mounting a Tec 7 Vaporizer on a Selectatec 7 Compatibility Block is not recommended.

⚠ WARNING

Do not lift or support the vaporizer by holding the control dial. Handle the vaporizer with care at all times.

Before mounting a vaporizer onto the Selectatec Series manifold, ensure that each manifold port valve O-ring is intact and that there is no foreign matter around the mating surfaces. A damaged O-ring and/or foreign matter around the mating surfaces can cause leaks.

Earlier versions of the Selectatec Series Mounted Manifold that provide mounting positions for three vaporizers require that if only two vaporizers are fitted, then the center position must be occupied. If the center position is not occupied, the interlock that helps ensure that only one vaporizer at a time can be turned ON is ineffective.

Do not use a vaporizer if the liquid level decreases below the minimum level.

Before using a vaporizer allow it to attain the ambient temperature of the location in which it has to be used.

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Mounting the vaporizer

Step 1

Set the dial to \bigcirc .



Figure 3-1 • Setting the concentration dial

Step 2

Unlock the locking lever.

- Turn the lever counter-clockwise.
- Make sure the lever releases.

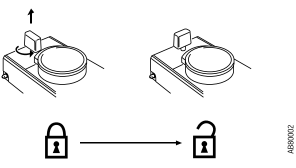
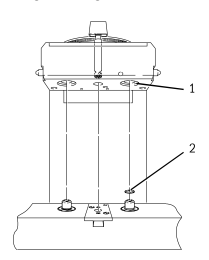


Figure 3-2 • Unlocking the locking lever

Step 3

Prepare the manifold.

- Remove any plugs fitted to the vaporizer interlock block ports.
- Verify that each manifold port valve O-ring is intact. If necessary, remove the existing O-rings and fit one new O-ring to each port valve, as described in the relevant anesthesia system User's Reference Manual. Replacement O-rings are supplied with each vaporizer.



1. Vaporizer Interlock Block Port - ensure plugs removed

2. Replace Manifold Port Valve O-ring, if necessary

Figure 3-3 ■ Readying the manifold

3-2

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Step 4

Install the vaporizer onto the manifold.

- Hold the main body of the vaporizer in an upright position with both hands.
- Lower the vaporizer onto the manifold, ensuring that the vaporizer interlock block ports engage correctly with the manifold port valves.

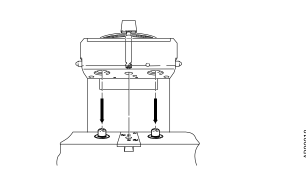


Figure 3-4 • Installing the vaporizer

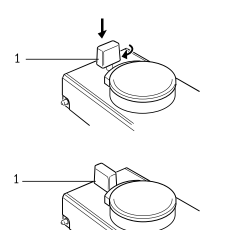
△ CAUTION

Push the locking lever all the way down before turning it. The mechanism can be damaged if an attempt is made to turn the lever before it is pushed all the way down.

Step 5

Lock the vaporizer onto the manifold.

- Push the locking lever all the way down.
- Turn it clockwise to the locked position to lock the vaporizer onto the manifold.



1. Locking lever

Figure 3-5 • Locking the vaporizer onto a manifold

Step 6

Ensure that the vaporizer is correctly mounted (see instructions on the next page).

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Checking the vaporizer for correct mounting

⚠ WARNING

To help ensure correct operation, do not use a vaporizer that is either visibly out of line on the manifold or that can be lifted off the manifold when the locking lever is in the locked position.

If more than one vaporizer is fitted, visually check to make sure that the tops of the vaporizers are level. If the vaporizer is visibly out of line, perform steps 2 and 3 as described in *Removing the vaporizer from a manifold* and remount it correctly.

When the vaporizer appears to be level and the locking lever is in the locked position, attempt to lift the vaporizer straight up from the manifold. If the vaporizer can be lifted off the manifold, it is not correctly mounted. Remount the vaporizer (see *Vaporizer mounting procedure*).

Verify that the interlock rods are in alignment by making sure that only one vaporizer at a time can be turned ON.

Check the anesthesia system for leaks in accordance with the relevant User's Reference Manual with the vaporizer dial turned to 0% and then repeat the check with the vaporizer dial turned to ∇ .

Removing the vaporizer from a manifold

Step 1

Set the dial to ∇ .

If the dial is not completely turned to the position the vaporizer cannot be released from the manifold.

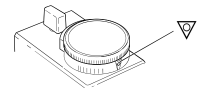


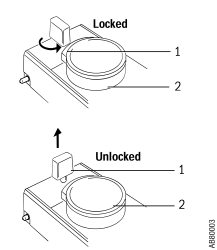
Figure 3-6 • Setting the dial

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Step 2

Unlock the locking lever.

- Turn the locking lever counterclockwise.
- Release the locking lever and check that the locking lever springs up to the unlocked position to release the vaporizer from the manifold.



- 1. Locking lever
- 2. Dial

Figure 3-7 • Unlocking the locking lever

Step 3

Carefully lift the vaporizer up from the manifold.

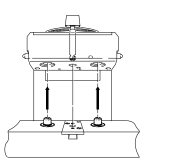


Figure 3-8 • Lifting the vaporizer

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3-6

4 Operating Instructions

Setting the dial

⚠ WARNING

High percent dial settings combined with low gas flows may lead to hypoxic mixtures in the breathing circuit. Datex-Ohmeda strongly recommends the use of oxygen monitoring.

The dial release must be operated to turn the dial from the ∇ setting.

Do not turn the dial if the vaporizer is not properly locked onto the manifold.

Important

European Standard EN 740 - Anesthetic Workstations and Their Modules requires that an appropriate gas monitor is used to monitor the concentration of anesthetic agent vapor in the inspiratory gas when the vaporizer is in operation in order to provide protection against hazardous output in the event of a device malfunction.

Datex-Ohmeda strongly recommends the use of anesthesia gas monitoring with this equipment. Refer to local standards for mandatory monitoring.

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Step 1

Press the dial release and turn the dial in a counter-clockwise direction from the setting.

Note that it is not possible to turn on the vaporizer if an adjacent Tec series, (except Tec 3) is turned on.

Step 2

The vaporizer should not be used between \bigcirc and the first graduation mark.

To avoid inadvertent delivery of small concentrations, turn the control dial to when the vaporizer is not in use.

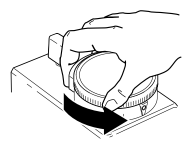


Figure 4-1 • Releasing the dial

4-2

Filling and draining the vaporizer

⚠ WARNING

Do not fill the vaporizer with any agent other than the agent specified on the front label. The vaporizer is designed for that agent only. If any substance other than that specified is used, patient injury could occur.

Only fill the vaporizer when it is in an upright position. Failure to do so may result in the vaporizer being overfilled.

To avoid explosive hazards, flammable anesthetic agents such as Ether and Cyclopropane must not be used in or with this vaporizer. Only anesthetic agents that comply with the requirements for non-flammable anesthetic agents in the IEC 60601-2-13 Standard, *Particular Requirements for the Safety of Anesthesia Machines*, are suitable for use in the presence of this vaporizer.

As this vaporizer is not suitable for use with flammable anesthetic agents such as Ether or Cyclopropane, the use of antistatic breathing tubes and face masks is not necessary. The use of antistatic or electrically conductive breathing tubes when using high frequency electric surgery equipment may cause burns and is therefore not recommended in any application of this vaporizer.

Do not fill the vaporizer unless the control dial is in the ∇ position.

Do not turn the dial ON during filling or attempt to fill beyond the **T** mark.

Do not drain the agent into any container other than a properly marked drug container.

Ensure that the filler cap is tightened prior to use.

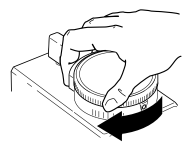
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When filling the Tec 7 Vaporizer, observe the following:

- Periodically check the agent level. The vaporizer should be refilled at appropriate intervals. The vaporizer is designed to function according to specification as long as there is agent visible above the mark.
- The vaporizer must be filled and used in an upright position. Small deviations from the upright position do not affect either the output or the safety of the vaporizer.
- Every two weeks, preferably when the agent level is low, drain the contents of the
 vaporizer into an appropriately marked container and discard the agent. Less
 frequent intervals may be used when the anesthetic agent does not contain
 additives or stabilizing agents, but the procedure must be performed at least
 once every year.
- The following steps should be taken for Halothane vaporizers:
 - Drain the vaporizer every two weeks
 - If Halothane is used infrequently the vaporizer should be drained after use.
 - The decomposition of halothane causes the release of halides, which may corrode metal components particularly in the presence of moisture. Also a preservative added to halothane by its manufacturers to impede decomposition can leave a residue, which may cause vaporizer components to stick.
- If the vaporizer is not upright, check the agent level more frequently to avoid a misleading impression of the amount of agent in the vaporizer.

Step 1

Turn the vaporizer dial clockwise to the position.



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Figure 4-2 • Turning the vaporizer OFF

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Filling procedure with funnel filler

⚠ WARNING

Before filling a vaporizer equipped with a funnel filler, turn the cap slowly to allow any pressure to gradually vent.

Ensure that the drain plug screw, located on the lower front of the vaporizer, is correctly tightened to help prevent loss of liquid agent.

Step 1

Remove the filler cap by turning it counter-clockwise. Ensure that the drain plug is closed by tightening it with the tool end of the filler cap.

Step 2

Verify that the anesthetic agent is the same as that specified on the vaporizer front label. Observe the agent level through the sight glass indicator on the side of the filler body. Pour the agent slowly into the filling port, as illustrated on Fig. 4-3, until the level reaches the mark. The level may decrease slightly as the wicks absorb the agent. To help prevent overfill, ensure the agent level is at or below the fill line.

Step 3

Replace and tighten the filler cap to minimize leaks.

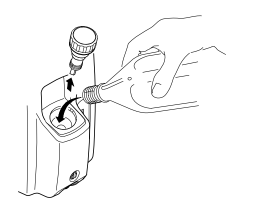


Figure 4-3 • Filling a vaporizer that incorporates a funnel filler



Figure 4-4 • Replacing the filler cap

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Draining procedure with funnel filler

The vaporizer must only be drained into a properly marked container.

▲ CAUTION

Do not allow the container to become completely full during draining procedures.

Step 1

Remove the filler cap. Insert the tool end of the cap into the drain plug below the filling port on the filler body as illustrated in Fig. 4-5.

Step 2

Position a properly marked container under the drain spout.

Step 3

Unscrew, but do not remove, the drain plug to allow the vaporizer contents to pour from the drain spout into the container.

Step 4

After draining is complete, tighten the drain plug to help minimize leaks.

Step 5

Replace and tighten the filler cap to help minimize leaks.

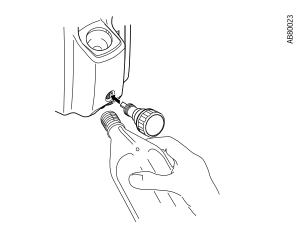


Figure 4-5 • Draining a vaporizer with a funnel filler



Figure 4-6 • Replacing the filler cap

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Filling procedure with Easy-Fil™

⚠ WARNING

Ensure that the drain plug screw, located on the lower front of the vaporizer, is correctly tightened to help prevent loss of liquid agent.

The filling system consists of three elements:

- the bottle collar
- the bottle adapter
- the filler block

The vaporizer must only be filled using the correct agent specific filling system.

Step 1

Align the notches on the bottle adapter to the bottle collar and tighten the adapter onto the agent bottle.

Step 2

Remove the filler cap. Align the bottle adapter keys with the index slots in the filler block as illustrated in Fig. 4-7. Insert the bottle nozzle into the filler block.

Step 3

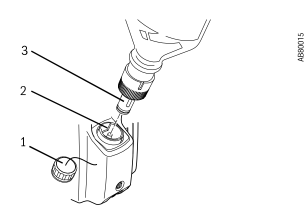
Press the agent bottle fully into the vaporizer filler block. Allow the liquid to flow into the vaporizer until the maximum level mark \blacksquare is reached. To help prevent overfill, ensure the agent level is at or below the fill line. Pay particular attention to the level in the sight glass and the air return bubbles flowing into the bottle.

Step 4

Release the bottle when the vaporizer is full and the continuous stream of bubbles ceases.

Step 5

Remove the bottle from the vaporizer filler. Replace the filler cap and the cap on the agent bottle. Ensure that the filler cap is tightened to help minimize leaks.



- 1. Filler cap
- 2. Index slot
- 3. Bottle adapter

Figure 4-7 • Filling a vaporizer with Easy-Fil

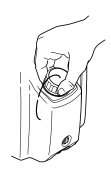


Figure 4-8 • Replacing the filler cap

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Draining procedure with Easy-Fil

⚠ CAUTION

Do not allow the container to become completely full during draining procedures.

The vaporizer must only be drained into a properly marked container.

Step 1

Remove the cap from the vaporizer filler.

Step 2

Place the empty container opening under the drain nozzle as shown in Fig. 4-9.

Step 3

Unscrew the drain plug with the tool attached to the filler cap. Drain the vaporizer until empty.

Step 4

After draining, tighten the drain plug to help minimize leaks.

Step 5

Replace and tighten the filler cap to minimize leaks.

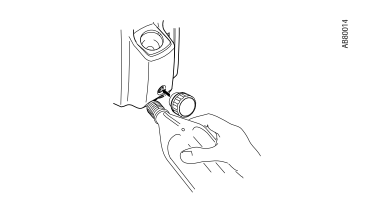


Figure 4-9 • Unscrewing the drain plug



Figure 4-10 • Replacing the filler cap

4-8

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Filling procedure with Quik-Fil™

⚠ WARNING

Ensure that the drain plug screw, located on the lower front of the vaporizer, is correctly tightened to help prevent loss of liquid agent.

Step 1

Remove the protective cap from the anesthetic agent bottle filler, checking that the bottle and filler mechanism are not damaged.

Step 2

Remove the filler cap. Insert the bottle nozzle into the filler block. Rotate the bottle to align the bottle filler nozzle keys with the index slots in the filler block as illustrated on Fig. 4-11.

Step 3

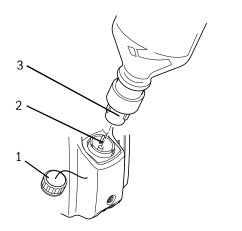
Press the agent bottle fully into the vaporizer filler block. Allow the liquid to flow into the vaporizer until the maximum level mark is reached. Pay particular attention to the level in the sight glass and the air return bubbles flowing into the bottle.

Step 4

Release the bottle when the vaporizer is full and the continuous stream of bubbles ceases. To help prevent overfill, ensure the agent level is at or below the fill line.

Step 5

Remove the bottle from the vaporizer filler. Replace the filler cap and the cap on the agent bottle. Ensure that the filler cap is tightened to help minimize leaks.



- 1. Filler cap
- 2. Index slot
- 3. Nozzle key

Figure 4-11 • Filling a vaporizer with a Quik-Fil



Figure 4-12 • Replacing the filler cap

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Draining procedure with Quik-Fil

⚠ CAUTION

Do not allow the container to become completely full during draining procedures.

The vaporizer must only be drained into a properly marked container.

Step 1

Remove the cap from the vaporizer filler.

Step 2

Place the empty container under the drain nozzle as shown in Fig. 4-13. Screw bottle onto drain spout.

Step 3

Unscrew the drain plug with the tool attached to the filler cap. Drain the vaporizer until empty.

Step 4

After draining, tighten the drain plug to help minimize leaks.

Step 5

Replace and tighten the filler cap to minimize leaks.

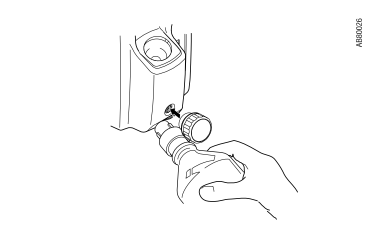


Figure 4-13 • Unscrewing the drain plug



Figure 4-14 • Replacing the filler cap

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5 Maintenance

User maintenance

⚠ WARNING

Do not modify, tamper with, or disassemble the vaporizer. Doing so can damage the unit and alter the graduation accuracy.

Maintenance intervals

Prior to performing any maintenance procedures or returning to a service center for repairs, clean and disinfect the vaporizer.

Every two weeks: When the agent is low, drain the contents of the vaporizer into an appropriately marked container and discard the agent. For Halothane vaporizers check the output of anesthetic agent periodically with an agent monitor. See note below.

12 months

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system by qualified personnel.

Inspect and perform output concentration check.

Note

The decomposition of Halothane causes the release of halides, which may corrode metal components particularly in the presence of moisture. Also a preservative added to Halothane by its manufacturers to impede decomposition can leave a residue, which may cause vaporizer components to stick. If Halothane is used infrequently the vaporizer should be drained after use.

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Cleaning

⚠ WARNING

Do not put water or any other solvent into a vaporizer. A vaporizer should be filled with the specified anesthetic agent only.

Do not immerse the vaporizer in water or any other liquid.

Do not autoclave the vaporizer.

External cleaning

To clean external surfaces, use a moist cloth and neutral detergent (pH 7 to 10.5).

Never allow cleaning agents to accumulate either in the filler, the gas inlet and outlet ports or around the control dial.

Internal contamination

If the vaporizer is filled or partly filled with an incorrect volatile agent or other contaminant (such as water), proceed as follows:

- Remove the vaporizer from service immediately and label the vaporizer stating that it is contaminated, Discard all liquid.
- 2. Return the vaporizer to a Datex-Ohmeda Authorized Service Center stating that the vaporizer is contaminated and, if possible, the type of contaminant in the vaporizer.

Output concentration check

Connect the Tec 7 to an Anesthesia Machine.

- 1. Set the oxygen output of the anesthesia machine to a flow of 5 ± 0.5 L/min.
- 2. Ensure that the fresh gas output is connected to a gas scavenging system.
- 3. Measure the concentration at the fresh gas outlet, using an agent monitor which is calibrated to measure the specific agent.
- 4. Allow the readings to stabilize and check that the readings are within specified tolerances.
- 5. Document and maintain the test results, including the date, person performing the test, and serial number of the unit tested.

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The accuracy of the measuring equipment must be considered when obtaining the readings!

Sevoflurane Dial Setting	Min Vol%	Max Vol%
1%	.6	1.40
3%	2.55	3.45
5%	4.25	5.75
Enflurane, Halothane or Isoflurane Dial Setting	Min Vol%	Max Vol%
	Min Vol% .75	Max Vol% 1.25
Isoflurane Dial Setting		

Checking the calibration

The performance of most vaporizers that are in clinical use can be confirmed by observing patient signs and consumption of anesthetic agents. Some users may, however, wish to employ analyzers either as a routine procedure or as part of an investigation to determine whether any abnormalities of performance have developed.

To help to achieve the reliability and consistency of performance of the Tec 7 Vaporizer, Datex-Ohmeda uses closely specified test conditions, test methods and detailed protocol in conjunction with training, experience and quality auditing systems. For these reasons, the full program necessary to help to ensure that a vaporizer complies with Datex-Ohmeda specifications cannot practicably be carried out in a field situation.

The following points must be considered when any measurements are being carried out on vaporizers to assist in determining whether any abnormalities of performance have developed.

- To predict the concentration that the vaporizer can be expected to deliver, the detailed nominal performance data and the preceding comments must be taken into account.
- The method of test used must not be such that it bears little relation to normal conditions of clinical use.
- 3. Any sampling techniques used must be such as to ensure the following:
 - a. The sample is fully representative of the vaporizer output, which may not be a homogeneous mixture at the vaporizer outlet.
 - b. The absorption of agent by any connecting tubing is negligible.
- 4. If a number of vaporizers are being examined at the same time the probability of all of them being consistently in error is so remote as to be negligible and the cause of any apparent error probably lies in the test method employed.
- 5. Consistent and reproducible analytical techniques must be used.
- 6. If unexpected results are obtained, it is a wise precaution to repeat the observation because the vaporizer may be more reliable than the techniques used to observe its performance.

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- If unexpected results occur, it is also worthwhile checking for sources of error such as the flowmeter, leaks or absorption by adjacent components.
- 8. Full account must be taken of any extraneous effects on the analyzer that may arise from changes in the carrier gas composition.
- 9. If the anesthetic machine on which the vaporizer is fitted is left for a period of time with no gases flowing, sensitive analyzers may detect small concentrations of agent for a short time at the machine outlet after the gas flow is turned ON with the vaporizer turned to . This is a normal machine characteristic caused by residual vapor left in the machine from previous use.
- 10. When the vaporizer is turned from ∇ to 0% or above after a period out of use, an increased concentration may occur that rapidly stabilizes to the set concentration within approximately 10 seconds at 5 liters/minute flow.
- At the 0% setting it is normal for small steady concentrations to be observable on sensitive analyzers.

Analytical techniques

For field checking of the state of calibration, many techniques and analyzers are available. Datex-Ohmeda would not recommend any one technique or analyzer in preference to another, but the calibration and reliability of analyzers must be realistically considered and account must be taken of errors of use.

The following method of checking can be used where special equipment is not available or where a secondary check of analyzers is desired. The characteristics of the vaporizer are such that, if the vaporizer is satisfactory at one dial setting, it should be satisfactory at all other graduations.

- 1. Ensure that the vaporizer is at least half full and has been at an ambient temperature of $21 \pm 2^{\circ}$ C for at least three hours.
- With the vaporizer securely mounted, drain the vaporizer as detailed in Section 4 and, after draining, ensure that either the drain plug and the filler cap are both securely tightened or the port valve is fully closed and the locking clamp is in the up position, as appropriate.
- 3. Check that the control dial is turned to ∇ and then carefully and quickly pour a measured 50 milliliters of agent into the vaporizer without spilling.
- 4. Leave the vaporizer at a nominal temperature of $21 \pm 2^{\circ}$ C for one hour to help to ensure that the temperature has stabilized.
- 5. Set the flowrate to 5 liters/minute oxygen.
- 6. Turn the control dial to 2%, note the time and check that the flowrate is still 5 liters/minute. Readjust the flowrate as necessary.
- 7. Leave the vaporizer at this setting for 30 minutes. Periodically check and adjust the flowrate as necessary. Turn the vaporizer to and turn the oxygen OFF.

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8. Drain the vaporizer as detailed previously in Instruction 2 and measure the amount of liquid drained off. The amount of liquid consumed should be as follows:

Enfluratec 15.5 milliliters
Fluotec 13.5 milliliters
Isotec 15.5 milliliters
Sevotec 16.6 milliliters

Appropriate action must be taken to handle the exhaust gases and spillage.

Note

This method is designed to be a quick and easy check of vaporizer operation and, therefore, it is somewhat imprecise. However, it is unusual for the measured liquid consumption to vary by more than 25% of the values listed above.

Service Policy

Repairs and service procedures must be performed at a Datex-Ohmeda Authorized Service Center. Contact your Datex-Ohmeda Service Representative or Datex-Ohmeda Authorized Distributor for information on maintenance and shipping.

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6 Principle of Operation

Interlock mechanism

The vaporizer locking lever is interlocked with the vaporizer percentage control dial so that the control dial release, located at the rear of the dial, cannot be actuated until the vaporizer locking lever is in the locked position.

With the vaporizer locking lever in the locked position, the dial release can be pressed in toward the dial to operate the interlock mechanism, which allows the manifold port valves to open, prevents an adjacent vaporizer from being turned on, and allows the vaporizer to operate.

Turning the control dial to \bigvee automatically reverses the operating sequence, which allows the dial release to move out to lock the dial in the \bigvee position, closes the manifold port valves and vents the vaporizer gas connecting ports, and allows an adjacent vaporizer to be turned ON.

Turning the locking lever to the unlocked position releases the vaporizer allowing it to be removed from the manifold.

Delivery of gas/agent vapor

Overview

The output concentration of the Tec 7 Vaporizer is regulated by the 'variable flow-split' method described in the following text and shown in figures 6-1 and 6-2.

A total flow of fresh gas from upstream flowmeters enters the vaporizer from the flowmeter where it is immediately split into two streams. One stream flows into the fresh gas bypass circuit and the other stream flows through the vaporizing chamber where it is enriched with the vapor of the liquid anesthetic agent.

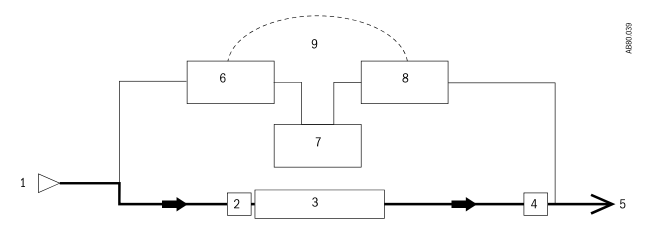
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Bypass circuit

The bypass circuit includes the gas transfer manifold and also a thermostat assembly that is located at the base of the vaporizer.

The fresh gas flows through the bypass circuit vertically downwards across the sump base through the thermostat and back up the gas transfer manifold to the common gas outlet as shown in Figure 6-1.

The thermostat deflects according to its temperature to control the resistance offered to the flow of gas through it. This deflection varies the relative proportions of gas flowing through the bypass and vaporizing chamber circuits.



- 1. Gas from flowmeter
- 2. Sump base
- 3. Thermostat
- 4. Gas transfer manifold
- 5. To common gas outlet
- 6. Shut-off open
- 7. Vaporizing chamber
- 8. Flow control (vapor channel)
- 9. Rotary valve

Figure 6-1 • Bypass circuit

Vaporizing chamber circuit

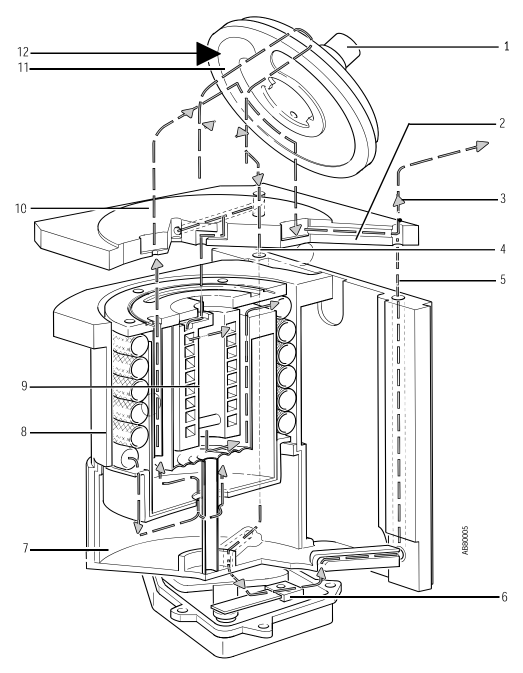
The fresh gas flow through the vaporizing chamber, as shown on Figure 6-2, flows from the flowmeter across the sump cover where it is diverted through the central cavity of the rotary valve and back through the Intermittent Positive Pressure Ventilation (IPPV) compensating assembly.

Gas now flows from the IPPV assembly down through the tubular wick assembly where it picks up anesthetic vapor and then flows across the base of the vaporizing chamber above the liquid agent.

From the base of the vaporizing chamber the gas/agent mixture flows through the sump cover to the proportional radial drug control groove of the rotary valve and then back into the sump cover where it combines with the fresh gas from the bypass circuit.

The combined total flow then flows out from the vaporizer and via the Selectatec circuitry to the anesthesia gas delivery system.

6-2



- 1. Rotary valve
- 2. Enriched fresh gas out
- 3. Combined fresh gas and enriched gas out
- 4. Fresh gas bypass
- 5. Fresh gas out
- 6. Thermostat
- 7. Vaporizing chamber
- 8. Wick assembly
- 9. IPPV compensating assembly
- 10. Sump cover
- 11. Vapor control channel
- 12. Shown in ON position

Figure 6-2 • Vaporizer schematic

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6-4

7 Specifications

Note All specifications are nominal and subject to change without notice.

Calibration

Check the calibration certificate that is included with your Tec 7 Vaporizer.

Vaporizers are calibrated at 21° C using an oxygen carrier gas at a flow of 5 liters/minute and they are temperature, flow and pressure compensated within the specified operating range.

⚠ WARNING

The Tec 7 Vaporizer can only be calibrated at a Datex-Ohmeda Authorized Service Center.

Performance

Accuracy at 5 liters/ min O_2 21 ± 2 °C	Vaporizer 5%: \pm 0.25% of delivered agent or \pm 15% dial setting (whichever is greater) Vaporizer 8%: \pm 0.4% of delivered agent or \pm 15% dial setting (whichever is greater)	
Liquid capacity	To fully charge a vaporizer with dry wicks: 300 ml (nominal) Retained by wick system: 75 ml (nominal) To fill from minimum to maximum mark: 5% Vaporizer: 170 ml (nominal) 8% Vaporizer (Sevotec): 137 ml (nominal)	
Flow resistance at 5 liters/minute of O_2 at 21 ± 2 °C	10 - 15 cm H ₂ O with Vaporizer setting ON +0%	
Operating temperature range	18 °C to 35 °C (64 °F to 95°F)	
Storage temperature range	-40 °C to 65 °C (-40 °F to 149°F) Note: Protect the vaporizer packaging from condensation.	

Weight and dimensions

Weight	7 kg	15 l b 6 oz (empty)
Depth	210 mm	8.25 inches
Width	110 mm	4.375 inches
Height	250 mm	9.875 inches (in the unlocked position)

Flow characteristics

Isotec 5%

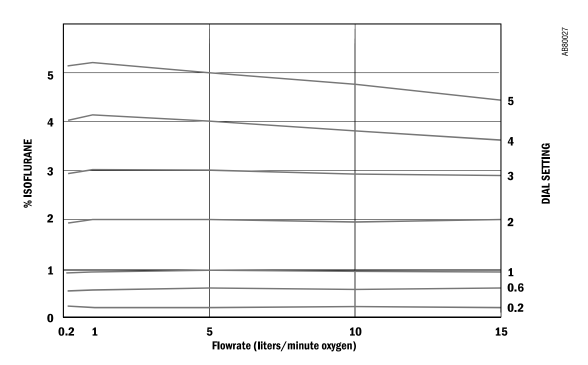


Figure 7-1 • Effect of Flowrate at 21 ± 2° C with oxygen flowing

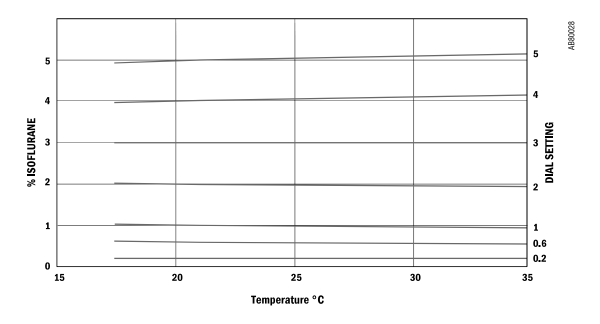


Figure 7-2 • Effect of temperature at 5 liters/minute with oxygen flow

Fluotec 5%

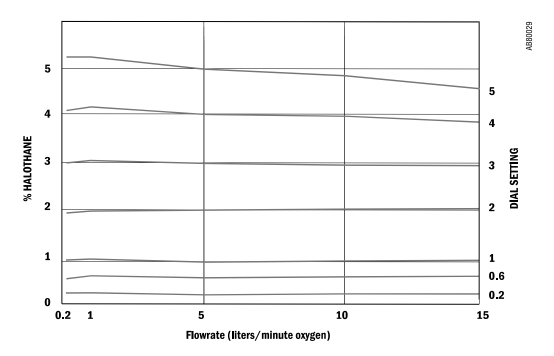


Figure 7-3 • Effect of Flowrate at 21 ± 2° C with oxygen flowing

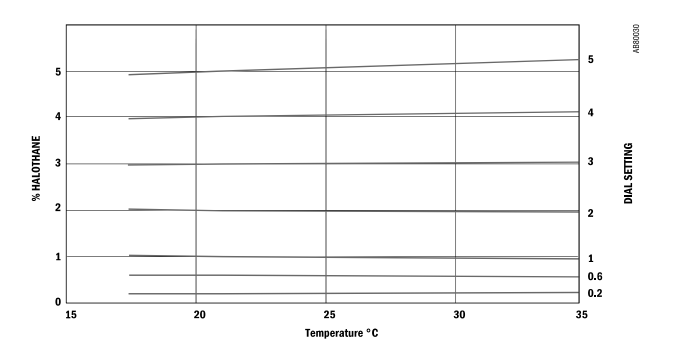


Figure 7-4 • Effect of temperature at 5 liters/minute with oxygen flow

Sevotec 5%

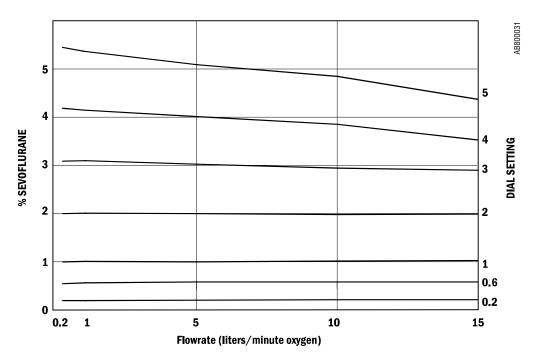


Figure 7-5 • Effect of Flowrate at 21 ± 2° C with oxygen flowing

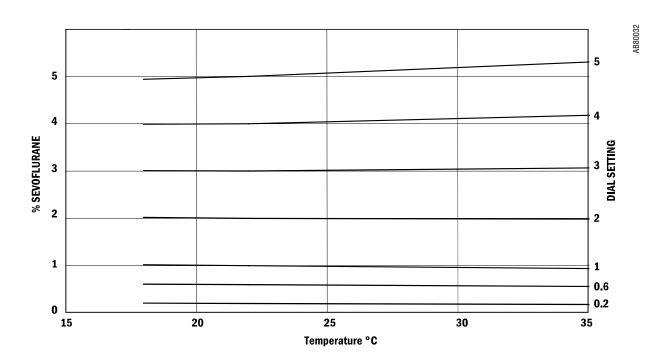


Figure 7-6 • Effect of temperature at 5 liters/minute with oxygen flow

Enfluratec 5%

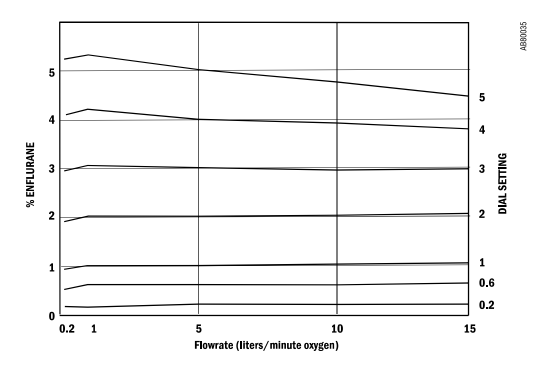


Figure 7-7 ■ Effect of Flowrate at 21 ± 2° C with oxygen flowing

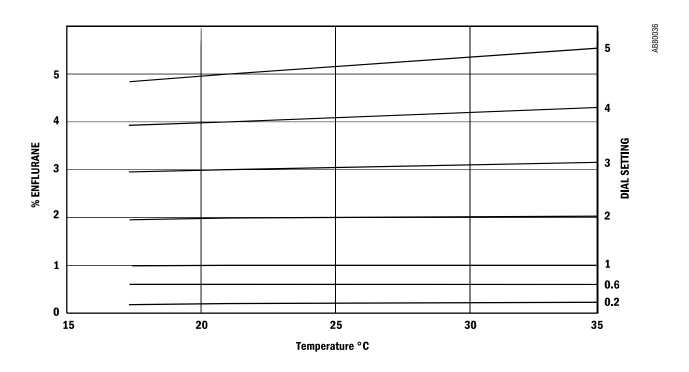


Figure 7-8 • Effect of temperature at 5 liters/minute with oxygen flow

Sevotec 8%

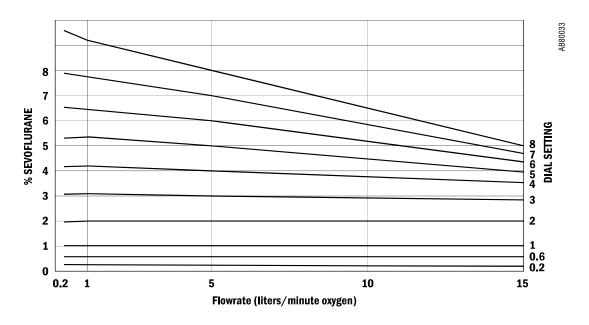


Figure 7-9 • Effect of Flowrate at 21 ± 2° C with oxygen flowing

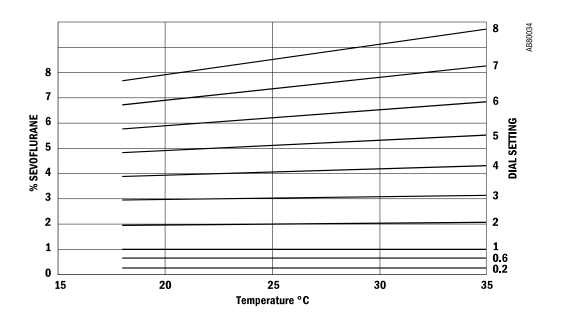


Figure 7-10 • Effect of temperature at 5 liters/minute with oxygen flow

Effects of variables

Anesthetic agent consumption

Isoflurane, Halothane and Enflurane

The rate of consumption of anesthetic agent depends primarily on flowrate and vapor output concentration. As an approximate working figure, 1 milliliter of liquid anesthetic is required to provide 200 milliliters of vapor.

The approximate hourly consumption of an esthetic agents can be expressed as 3×7 , where 7×7 represents the setting of the vaporizer output percentage and F represents the input flowrate in liters/minute.

Example: If a vaporizer is set to deliver 2% at 6 liters/minute total gas

input flowrate then the approximate rate of consumption

 $= 3 \times 2 \times 6 = 36 \text{ ml/hour.}$

The figures are approximate and are intended for general guidance only.

Sevoflurane

The rate of consumption of anesthetic agent depends primarily on flowrate and vapor output concentration. As an approximate working figure, 1 milliliter of liquid anesthetic is required to provide 200 milliliters of vapor.

The approximate hourly consumption of anesthetic agents can be expressed as $3.3 \, x \, \% \, x \, F$, where % represents the setting of the vaporizer output percentage and F represents the input flowrate in liters/minute.

Example: If a vaporizer is set to deliver 2% at 6 liters/minute total gas input flowrate then the approximate rate of consumption

 $= 3.3 \times 2 \times 6 = 39.6 \text{ ml/hour.}$

The figures are approximate and are intended for general guidance only.

Barometric pressure

The Tec 7 Vaporizer is calibrated in percent v/v at 760 mmHg. If the ambient pressure changes the % v/v changes, so that at an ambient pressure P mmHg the delivered percentage (D% v/v) is calculated as follows:

$$D = \frac{\% \times 760}{P}$$
 where % is the nominal setting of the vaporizer.

To obtain a consistent depth of an esthesia when gross changes of barometric pressure occur, the % v/v must be changed in inverse proportion to the barometric pressure.

The vaporizer automatically makes this % v/v change and for practical clinical purposes the effects of barometric pressure can be ignored.

Ambient temperature

The effects of variation in temperature are normally negligible at commonly used combinations of dial setting and ambient temperature.

If the vaporizer temperature is above the range shown on the performance curves, the vaporizer output may be unpredictably high, particularly if the temperature of the agent approaches the agent boiling point specified by the agent manufacturer.

If the vaporizer temperature is below the range shown on the performance curves, the vaporizer output may be lower than expected.

To help avoid inaccuracies due to extreme temperatures, before using the vaporizer it must be allowed to attain a temperature within the range shown on the performance curves.

Back pressure

⚠ WARNING

Pressures in excess of 400 mmHg may overcome the internal pressure balance and cause a variation in output.

Steady back pressure

The vaporizer cannot distinguish between pressures at the outlet due to barometric pressure and pressures in excess of barometric due to steady back pressures applied by downstream components. The equation given in the section Barometic Pressure therefore applies with the term P now being the absolute pressure at the outlet, that is, barometric pressure plus back pressure. Steady back pressure reduces the % v/v.

Currently, it is unlikely that the steady back pressure imposed by commonly used downstream components, other than some ventilators, exceeds 30 mmHg at commonly used flowrates. Back pressures as high as 30 mmHg would reduce the delivered % v/v, at 760 mmHg barometric pressure, to the following:

 $\frac{700}{790}$ = 0.96 of what would otherwise be expected.

Under normal clinical circumstances effects of this magnitude can be ignored.

Fluctuating back pressure

Fluctuating back pressure may be imposed on the vaporizer by downstream components and/or assisted or controlled ventilation to the patient. These fluctuating back pressures can affect the vaporizer and increase the concentration by intermittently altering the pressures, and consequently the flow distribution, within the vaporizer.

The greatest effects are observed at combinations of very low flowrates and low dial setting with large and rapid pressure fluctuations. The effects become progressively less important as the dial setting and flowrate increase and the magnitude and rate of cycling of the pressure fluctuations decrease.

Carrier gas composition

Small output decreases can occur when the carrier gas composition is changed from 100% oxygen.

When either air or nitrous oxide is employed as the carrier gas, the output is lowered compared to the output when oxygen is the carrier gas. This effect is the greatest (up to 20% of setting) at low flows when nitrous oxide is employed, but using nitrous oxide reduces the required inspired concentrations of volatile agent that can, depending upon the proportion, mitigate the decreases in output from the vaporizer.

⚠ WARNING

Only operate the vaporizer with dry medical gases.

Time out of service

If the anesthesia machine on which the vaporizer is fitted is left for a period of time with no gases flowing, small concentrations may be detected at the machine outlet immediately after the gas flow is turned ON. This is a normal machine characteristic and is caused by residual vapor left in the machine from previous use.

When the vaporizer is turned from the setting after a period out of use, a brief high concentration may occur that rapidly stabilizes to the set concentration within approximately 10 seconds at 5 liters/minute.

These phenomena are normal characteristics of vaporizers. In use the volume of vapor involved is small compared to the volume of the breathing circuit.

Effects of variables

Ambient temperature, input flowrate and duration of use can affect delivered concentration, particularly when the vaporizers are used at extremes of the usual clinical range.

Note

Use of the vaporizer at high gas flows and high dial concentrations may affect the accuracy of delivered concentrations. Refer to Performance Curves in this chapter for full information.

The valve design and temperature compensation system of Tec 7 Vaporizers reduces the effects to levels such that, under most clinical conditions, their effect on vaporizer performance is not clinically significant.

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Warranty

This Product is sold by Datex-Ohmeda under the warranties set forth in the following paragraphs. Such warranties are extended only with respect to the purchase of this Product directly from Datex-Ohmeda or Datex-Ohmeda's Authorized Dealers as new merchandise and are extended to the Buyer thereof, other than for the purpose of resale.

For a period of 36 months from the date of original delivery to Buyer or to Buyer's order, but in no event for a period of more than three years from the date of original delivery by Datex-Ohmeda to a Datex-Ohmeda Authorized Dealer, this Product, other than its expendable parts, is warranted against functional defects in materials and workmanship and to conform to the description of the Product contained in this User's Reference Manual and accompanying labels and/or inserts, provided that the same is properly operated under the conditions of normal use, that regular periodic maintenance is performed and that replacements and repairs are made in accordance with the instructions provided. This same warranty is made for a period of thirty (30) days with respect to expendable parts. The foregoing warranties shall not apply if the Product has been repaired other than by Datex-Ohmeda or in accordance with written instructions provided by Datex-Ohmeda, or altered by anyone other than Datex-Ohmeda, or if the Product has been subject to abuse, misuse, negligence, or accident.

Datex-Ohmeda's sole and exclusive obligation and Buyer's sole and exclusive remedy under the above warranties is limited to repairing or replacing, free of charge, at Datex-Ohmeda's option, a Product, which is telephonically reported to the nearest Datex-Ohmeda Customer Support Center and which, if so advised by Datex-Ohmeda, is thereafter returned with a statement of the observed deficiency, not later than seven (7) days after the expiration date of the applicable warranty, during normal business hours, transportation charges prepaid, and which, upon Datex-Ohmeda's examination, is found not to conform with above warranties. Datex-Ohmeda shall not be otherwise liable for any damages including but not limited to incidental damages, consequential damages, or special damages.

There are no express or implied warranties which extend beyond the warranties hereinabove set forth. Datex-Ohmeda makes no warranty of merchantability or fitness for a particular purpose with respect to the product or parts thereof.