INFOV.A.C.™
THERAPY SYSTEM
USER MANUAL

Rx Only
Indications, Contraindications, Warnings, Precautions and other Safety Information are contained in the V.A.C.* Therapy System Safety Information Sheet. This information sheet is included with the therapy unit and also included in V.A.C.* Dressing cartons. Please consult the V.A.C.* Therapy System’s User Manual and the Safety Information Sheet before applying V.A.C.* Therapy. If there are questions, or if this information sheet is missing, immediately contact your local KCI representative.

Additional product information can be found at www.acelity.com (USA) or www.kci-medical.com (outside the USA).

As with all prescription medical devices, failure to follow product instructions or adjusting settings and performing therapy applications without the express direction and/or supervision of your trained clinical caregiver may lead to improper product performance and the potential for serious or fatal injury. For medical questions, please consult a physician. In case of medical emergency, immediately contact your local emergency services provider.

CAUTION: Federal law (US) restricts this device to sale or rental by or on the order of a physician.

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Warnings And Important Information For Users

In order for KCI products to perform properly, KCI recommends the following conditions. Failure to comply with these conditions will void any applicable warranties.

- Use this product only in accordance with this manual and applicable product labeling.
- Assembly, operations, extensions, re-adjustments, modifications, technical maintenance or repairs must be performed by qualified personnel authorized by KCI. For these authorized personnel, KCI will make available upon request circuit diagrams, component parts lists, etc. as required for repairs.
- Ensure the electrical installation of the room complies with the appropriate national electrical wiring standards. To avoid the risk of electrical shock, this product must be connected to a grounded power receptacle.
- Do not operate this product if it has a damaged power cord, power supply or plug. If these components are worn or damaged, contact KCI.
- Do not drop or insert any object into any opening or tubing of this product.
- Do not connect this product or its components to devices not recommended by KCI.
- Use only V.A.C.* Dressings and disposables with this product.
- Keep this product away from heated surfaces.
- Although this product conforms to the intent of the standard IEC 60601-1-2 in relation to Electromagnetic Compatibility, electrical equipment may produce interference. If interference is suspected, separate the equipment and contact KCI.
- Avoid spilling fluids on any part of this product.

⚠️ Fluids remaining on the electronic controls can cause corrosion that may cause the electronic components to fail. Component failures may cause the unit to operate erratically, possibly producing potential hazards to patient and staff. If spills do occur, unplug the unit immediately and clean with an absorbent cloth. Ensure there is no moisture in or near the power connection and power supply components before reconnecting power. If the product does not work properly, contact KCI.

- Do not use this product while bathing/showering or where it can fall or be pulled into a tub, shower or sink.
- Do not reach for a product that has fallen into water. Unplug the unit immediately if plugged into electrical source. Disconnect the unit from dressing and contact KCI.
- Do not use this product in the presence of a flammable anesthetic mixture with air, oxygen, or nitrous oxide, or an oxygen enriched environment.
- Do not take this product into an MR environment. This product is MR Unsafe.

Notice

This product has been configured from the manufacturer to meet specific voltage requirements. Refer to the Product Information Label for specific voltage.
Introduction

V.A.C.® (Vacuum Assisted Closure®) Therapy is a system that uses controlled continuous or intermittent negative pressure (vacuum) to create an environment that promotes wound healing by secondary or tertiary (delayed primary) intention by:

- preparing the wound bed for closure
- reducing edema
- promoting granulation tissue formation and perfusion
- removing exudate and infectious material

The INFOV.A.C.™ Therapy System provides Negative Pressure Wound Therapy (NPWT) and Therapeutic Regulated Acute Care (SENSAT.R.A.C.™) for use on a variety of chronic and acute wound types. This advanced wound healing technology, coupled with microprocessor-controlled therapy units, and first-class technical back-up, can be readily integrated into the clinician’s wound healing practice, helping to optimize patient care and manage costs.

Protection Against Hazards

Discard all disposable items (all tubing, connectors, clamps, used canister, used dressings, etc.) in accordance with local medical waste disposal regulations.

Standard Precautions should be used when handling any body fluids or waste. Properly dispose of all parts according to institutional procedures, as well as local, state and federal regulations.

The INFOV.A.C.™ Therapy Unit and power supply should be returned to KCI for disposal at the end of their operational life.

Only the power supply provided with the INFOV.A.C.™ Therapy Unit should be used to power the device or to recharge the battery. Using any other power supply may damage the INFOV.A.C.™ Therapy Unit.

The battery in the INFOV.A.C.™ Therapy Unit is not a user serviceable part. If battery problems are suspected, the system should be returned to KCI for servicing.

Power cords and tubing may present a tripping hazard. Ensure that all cords and tubing are out of areas where people may walk.
Attaching the INFOV.A.C.™ Therapy Unit to Other Devices

The INFOV.A.C.™ Therapy Unit may be attached to the footboard of a hospital bed, most wheelchairs or to an IV pole (see Preparation for Use section). If required, it may be placed on a solid, level surface where it does not cause an obstruction. Ensure unit is placed below the height of the wound and where cables and tubes cannot be caught on passing objects.

The INFOV.A.C.™ Therapy Unit is not intended to be carried by ambulatory patients. Consult your physician and contact your local KCI representative for V.A.C.® Therapy units designed for ambulatory patient use.

Devices that can be used with the INFOV.A.C.™ Therapy Unit:

- Memory cards
- Infrared devices (available on some units)
- USB devices
INFOV.A.C.™ Therapy Unit - Parts Identification

- Power On/Off Button
- Canister Release Button
- Touch Screen User Interface
- INFOV.A.C.™ 500mL Canister
- Memory Card Slot
- Clinician Use Only
- Stylus
- Data Door
- Infrared (IR) Data Port
  - Available on some units, Clinician Use Only
- USB Data Port
  - Clinician Use Only
- Carry Handle
- Battery Charging Indicator Light
- Hanger Knob
- Hanger Arm
- Power Connection
- INFOV.A.C.™ Power Supply
Preparation for Use

Before preparing the INFOV.A.C.™ Therapy Unit for use, inspect the unit for any damage or contamination. Refer to the Care and Cleaning section of this manual for more information.

Charging the Battery

The INFOV.A.C.™ Therapy Unit comes with its own power supply and rechargeable battery. The battery is not user accessible or serviceable. The power supply has a two-part cord; one that plugs into an AC wall outlet and one that plugs into the INFOV.A.C.™ Therapy Unit. The AC power cord may have different wall plug configurations depending on country requirements.

> Use only the power supply provided with the INFOV.A.C.™ Therapy Unit to power the device or to recharge the battery. Using any other power supply may damage the INFOV.A.C.™ Therapy Unit.

> To isolate the therapy unit from supply mains, unplug the AC power from the wall outlet. Do not block access to the plug or wall outlet.

Power cords and tubing may present a tripping hazard. Ensure that all cords and tubing are out of areas where people may walk.

1. Plug the AC power cord into the power supply.

2. Plug the AC power cord into an AC wall outlet.

3. Locate the white arrow on the INFOV.A.C.™ power supply charging cord connector.

4. Plug the INFOV.A.C.™ power supply charging cord connector into the power connection on the back of the INFOV.A.C.™ Therapy Unit with the white arrow facing up.

It should take approximately four hours to fully recharge the battery. The system can be used while the battery is recharging.

To maximize battery life, keep the unit plugged in.
Battery Charging Indicator Light

When the INFOV.A.C.™ power supply is correctly plugged into the INFOV.A.C.™ Therapy Unit, the battery charging indicator light on the back of the unit will glow amber as the battery is charging. When the battery is fully charged the battery charging indicator light will glow green.

The battery level is shown on the bottom of the touch screen user interface:

- ![Battery Full](image)
  - Fully Charged
- ![Battery In Use](image)
  - In Use
- ![Battery Low](image)
  - Battery Low, charge battery soon
Attaching the INFOV.A.C.™ Therapy Unit to an IV Pole

1. Ensure the hanger arm is in the horizontal position.

2. Hold the INFOV.A.C.™ Therapy Unit by the carry handle, grip the hanger knob and pull the hanger arm out.

3. Place the hanger around the IV pole ensuring that the pole is in the vertical rubber groove on the rear of the INFOV.A.C.™ Therapy Unit. Allow the hanger to close pulling the therapy unit onto the pole.

4. Turn the hanger knob to lock the hanger arm in place. The lock symbol on the hanger knob will align with the arrow on the therapy unit when the mechanism is locked. An audible click also indicates that the mechanism is locked.

*The hanger knob can be turned past the lock position to further secure the INFOV.A.C.™ Therapy Unit. Hand tighten only; excessive force or tools should not be used.*
Attaching the INFOV.A.C.™ Therapy Unit to a Bed Footboard

1. Ensure the hanger arm is in the vertical position.

2. Hold the INFOV.A.C.™ Therapy Unit by the carry handle, grip the hanger knob and pull the hanger arm out.

3. Place the hanger over the footboard. Allow the hanger to close pulling the therapy unit onto the footboard.

4. Turn the hanger knob to lock the hanger arm in place. The lock symbol on the hanger knob will align with the arrow on the therapy unit when the mechanism is locked. An audible click also indicates that the mechanism is locked.

   The hanger knob can be turned past the lock position to further secure the INFOV.A.C.™ Therapy Unit. Hand tighten only; excessive force or tools should not be used.

Releasing the INFOV.A.C.™ Therapy Unit from an IV Pole or Bed

1. Hold the INFOV.A.C.™ Therapy Unit by the carry handle.

2. Reverse the attaching procedure.
Installing the 500mL and 1000mL Canisters

1. Press Power On/Off to turn the INFOV.A.C.™ Therapy Unit on.

2. Attach the canister:
   - Slide the canister into the end of the INFOV.A.C.™ Therapy Unit as shown below.
   - Push the canister firmly into place. An audible click signals the canister is correctly installed.

3. Connect the canister tubing to the dressing tubing:
   - Push the connectors together and twist until the locking tabs are fully engaged.

4. Open all tubing clamps as shown below.

   Closed Clamp

   Open Clamp
Changing the 500mL and 1000mL Canisters

A canister may be changed under routine conditions or under alarm conditions.

Under routine conditions the Canister Release button will NOT be flashing. When changing the canister under routine conditions do NOT turn power off to the INFOV.A.C.™ Therapy Unit.

Under alarm conditions the Canister Release button will be flashing. An alarm screen will be displayed and therapy will be off.

**WARNING:** According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.

1. Stop therapy by pressing the Therapy On/Off button on the touch screen user interface.

2. Close all tubing clamps as illustrated.

3. Disconnect the canister tubing from the dressing tubing:
   - Twist connectors until the locking tabs are fully disengaged and pull the connectors apart.
Changing the 500mL and 1000mL Canisters (cont)

4. Press the **Canister Release** button. The canister will eject but remain in the cradle.

5. Lift the canister out of the cradle.

6. Dispose of the canister according to institution and / or local environmental regulations.

**WARNING:** According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.
Installing the 300mL ACTIV.A.C.™ Canister

1. Attach the canister:
   - Push the canister firmly into place on the INFOV.A.C.™ Therapy Unit as illustrated below.

2. Connect the canister tubing to the dressing tubing:
   - Push the connectors together and twist until the locking tabs are fully engaged.

3. Open all tubing clamps as illustrated.

   ![Closed Clamp](image1)
   ![Open Clamp](image2)
Removing the 300mL ACTIV.A.C.™ Canister

When the Canister Release button is pressed, the 300mL canister is NOT held in place by the cradle of the INFOV.A.C.™ Therapy Unit. When removing the 300mL canister from the INFOV.A.C.™ Therapy Unit ensure that the canister is held FIRMLY before pressing the Canister Release button.

1. Hold the canister firmly.
2. Press the Canister Release button.

Operating Instructions

Before powering the therapy unit on, ensure that:

- The V.A.C.® Dressing is applied as described in the Application Instructions supplied with the V.A.C.® Dressing.
- The INFOV.A.C.™ Canister is attached as described in the Installing the Canisters sections of this manual.
- The INFOV.A.C.™ Therapy Unit is fitted to an IV pole or footboard or placed as described in the Preparation for Use section of this manual.

Powering the INFOV.A.C.™ Therapy Unit On / Off

Power On:
1. Connect the INFOV.A.C.™ Therapy Unit to the power supply as described in the Preparation for Use section of this manual.
2. Press and hold the Power On/Off button for approximately two seconds to turn the INFOV.A.C.™ Therapy Unit on. The Power On/Off button will illuminate and the touch screen user interface will activate.
   
   The INFOV.A.C.™ Therapy Unit start up process will briefly be displayed followed by a Warning Screen.

Power Off:
Press and hold the Power On/Off button for approximately two seconds to turn the INFOV.A.C.™ Therapy Unit off.

The Power On/Off button will flash for ten seconds after which the INFOV.A.C.™ Therapy Unit will automatically shut down.

WARNING: According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.
Operating the Touch Screen

The **touch sensitive** screen on the front of the unit displays information on current system operations and settings, and provides an interface for changing settings and operation.

The operation of the touch screen is detailed in the following pages.

The touch screen should ONLY be operated by finger or the supplied stylus (see [INFOV.A.C.™ Therapy Unit Parts Identification section](#)). Using pens or pointing devices will damage the screen and may affect the proper function of the unit.

Control Screens

The three levels of Control Screens are Patient Mode, Clinician Mode and Restricted Mode.

Patient Mode

The clinician may select Patient Mode to allow a patient to control certain elements of their therapy. This mode does not allow the patient to access or alter therapy settings.

The patient can activate Night Mode and the Screen Guard, access the Help screens, and respond to alarms.

Clinician Mode

The Clinician Mode screens allow a clinician to set and change therapy settings, view therapy history and access the Restricted Mode (see below).

Restricted Mode (Patient History)

The Restricted Mode screens allow access to patient wound images, image area trend charts and other patient health information that the caregiver has entered. From this mode the clinician can adjust the Leak Alarm sensitivity, and export therapy and patient histories. A secure password system protects the Restricted Mode from unauthorized access.

The secure password is created by the healthcare professional, not KCI. If a new password is created, all patient history associated with the previous password is deleted. All information is automatically deleted when therapy unit is returned to KCI.
Patient Mode Control Screens

Common Screen Control Buttons

- **Help Button**
  - Press to access Help Menu.

- **Screen Guard**
  - Press to activate Screen Guard feature to help prevent unintentional screen changes. This feature should be used when cleaning the touch screen. To release the Screen Guard, press 1 and then 2, which will appear on the screen when Screen Guard is active.

Navigation Buttons

- **Exit**
  - Press to leave the current screen.

- **Cancel**
  - Press to stop action in progress.

- **Next**
  - Press to go to the next screen.

- **Back**
  - Press to return to the previous screen.
Night Mode Function

The Night Mode function on the Patient Home Screen can be used to dim disrupting lights from the therapy unit during nighttime therapy use.

Press to activate the Night Mode function from the Patient Home Screen. A confirmation screen will be displayed before Night Mode is immediately activated.

When the Night Mode function is active, therapy continues as normal. The touch screen user interface will darken and the green Power On/Off button will dim.

The Night Mode function will deactivate in the event of an alarm.

During Night Mode, the screen will illuminate when touched and display the Patient Home Screen.

Should the touch screen user interface be touched while night mode is active, the screen will automatically darken after two minutes of inactivity.

Press to deactivate the Night Mode function.
Patient Mode Help Menu Screen

Press to access the Patient Mode Help Menu screen from any Patient Mode screen where this button appears.

Press to access Operating Instructions.

Press to access the About screen for information regarding the INFOV.A.C.™ Therapy Unit software.

Press to access Clinician Mode (press and hold for three seconds).

Press to access the Language screen to select the appropriate language.

Press to return to the Patient Home Screen.

Operating Instructions Screen

The Operating Instructions provide help in operating the INFOV.A.C.™ Therapy Unit.

Press to access the Operating Instructions screen from the Help Menu screen to view basic instructions for operation, cleaning and alarms.
Clinician Mode Control Screens

Common Screen Control Buttons

Press to access Clinician Mode Help Menu.

Press to activate Screen Guard feature to help prevent unintentional screen changes. This feature should be used when cleaning the touch screen. To release the Screen Guard, press 1 and then 2, which will appear on the screen when Screen Guard is active.

To Return to the Patient Home Screen

Press the Help button.

Press Patient Mode on the Help Menu screen.

Press and hold OK until the Patient Home Screen returns.
Therapy Start Screen

From the Clinician Home Screen press Therapy On/Off to access the Therapy Start screen.

A green bar graph indicates that the INFOV.A.C.™ Therapy System is operating normally.

Ensure that a new V.A.C.® Dressing has been applied and therapy settings have been selected per physician’s orders before starting therapy.

From this screen the clinician can use the SEAL CHECK™ Feature to view the integrity of the V.A.C.® Dressing, or use the Log tool to record a canister change or record the number of foam pieces used during a dressing change.

SEAL CHECK™ Feature Overview

The SEAL CHECK™ Feature is used to help find negative pressure leaks.

The SEAL CHECK™ Feature includes:

- an audible tone that changes frequency as the rate of the leak changes. Pressing the Seal Audio button will switch between the audible tone being on or off.
- a real time bar graph that gives a visual indication of the rate of the leak.

Access the SEAL CHECK™ Feature three different ways:

- When Therapy is started from the Clinician Home Screen, discussed on the following pages.
- When the SEAL CHECK™ Button is pressed on the Therapy screen.
- When the SEAL CHECK™ Button is pressed on the Leak Alarm screen after the INFOV.A.C.™ Therapy Unit detects a possible leak.

Patients only have access to the SEAL CHECK™ Feature through the Leak Alarm screen when the INFOV.A.C.™ Therapy Unit detects a possible leak.
How to Use the SEAL CHECK™ Feature (When Starting Therapy)

From the Clinician Home Screen press Therapy On/Off to access the Therapy Start screen.

An orange bar graph indicates a significant leak.

The SEAL CHECK™ Feature provides an audible tone and bar graph to assist in finding leaks. The frequency of the audible tone and the height of the bar graph will reflect the leak rate. The audible tone slows down and the bar graph decreases in height as the leak is found and corrected.

During initial dressing draw down, the bar graph should turn orange and then return to green if there are no significant leaks.

Most leaks occur in three specific areas.
- where the V.A.C.® Drape meets the skin.
- where the SENSAT.R.A.C.™ Pad is attached to the V.A.C.® Drape.
- at tubing connections.

Finding a Leak Using the SEAL CHECK™ Feature

1. Ensure connector between dressing tubing and canister tubing is properly locked.

2. Ensure the canister is securely installed on the therapy unit. If the canister is properly installed, the canister cannot be removed by gently pulling the canister directly away from the unit.

3. While therapy is on, apply gentle pressure and move your hand and fingers slowly around the edges of the drape and SENSAT.R.A.C.™ Pad. The bar graph will lower and the frequency of the audible tone (if Seal Audio is on) will decrease when the leak is found.

4. Refer to the Application Instructions provided with V.A.C.® Dressings for information on using excess V.A.C.® Drape material to seal the leak area.

5. When finished press Exit to return to the Clinician Home Screen.
Log Tool Overview

The Log tool can be used to track:

- the number of foam pieces used during a dressing change.
- canister changes.

Logged information is viewable and exportable on the Therapy History screens.

How to Use Log Tool (When Starting Therapy)

**Item to Log**

From the Clinician Home Screen press Therapy On/Off then Log to access the Item to Log screen.

Press to log the time and date of a canister replacement.

Press to log the number of foam pieces used in a dressing change.

Press to return to the Clinician Home Screen.

**Canister Replaced Screen**

From the Clinician Home Screen press Therapy On/Off then Log, then Canister to access the Canister Replaced screen.

Press to log that the canister has been replaced and return to the Item to Log screen. The current time and date will be recorded.

Press to return to the Item to Log screen without logging an entry.
From the Clinician Home Screen press Therapy On/Off then Log, then Dressing to access the Number of Foam Pieces screen.

Information displayed represents the last logged entry.

Use the + and - buttons to select the number of pieces of foam used during the current dressing change.

Press to log the number of foam pieces used and return to the Item to Log screen. The current time and date will be recorded.

Press to return to the Item to Log screen without logging an entry.

Always document the number of foam pieces used in patient chart and on the V.A.C.® Drape.

Logged information will appear in Therapy History as follows:

<table>
<thead>
<tr>
<th>dd/mmm/yy</th>
<th>Time</th>
<th>Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>11/Jan/10</td>
<td>15:54</td>
<td>Canister Changed</td>
</tr>
<tr>
<td>11/Jan/10</td>
<td>15:55</td>
<td>Dressing Changed, 4</td>
</tr>
</tbody>
</table>

The numeral after “Dressing Changed” is the number of foam pieces recorded on the above screen.
Therapy Screen

The Therapy screen enables clinicians to change settings, select wound type, check the seal and view the therapy or patient history.

From the Clinician Home Screen press Therapy to access the Therapy screen.

Press to manually set therapy.

Press to access the settings guide for preset therapy settings.

Press for assistance in finding leaks.

Press to view or export therapy history.

Press to view or export patient history from the Enter Access Code screen.

Settings Screen (Manual)

From the Clinician Home Screen press Therapy, then OK, then Settings to access the Settings screen.

Press to change pressure settings.

Press to change intensity.

Press to toggle between Continuous and Intermittent therapy.

Press to set Intermittent therapy times.
Pressure Screen

From the Clinician Home Screen press Therapy, then OK, then Settings then Pressure to access the Pressure screen.

Use the + and - buttons to change the desired pressure.

Pressure can be set from 25 to 200 mmHg in increments of 25 mmHg. Press Exit to go to the Confirm screen.

Default setting is: 125 mmHg (accuracy + / - 10 mmHg).

Intensity Control

From the Clinician Home Screen press Therapy, then OK, then Settings to access the Intensity button.

- Intensity is related to the time it takes to reach the target therapy level after the initiation of therapy.
- The lower the intensity setting, the longer it will take for the target therapy level to be reached.
- KCI recommends that new patients begin therapy at the lowest intensity setting as this allows for slower increase of negative pressure once the foam is compressed in the wound.
- The intensity can remain at the minimum setting throughout the entire length of treatment, if desired.

Press to change levels. Green crescent changes with each setting.

Default setting is: Low.
From the Clinician Home Screen press Therapy, then OK, then Settings, then Intermittent to access the Intermittent screen.

Use the + and - buttons to change the desired On and Off Time (in minutes).

Both On and Off Times can be set from 1 minute to 10 minutes in 1 minute increments.

Press Exit to go to the Confirm screen.

Changing Intermittent time intervals will take effect in the next cycle.

Default setting is: On Time = 5 minutes, Off Time = 2 minutes.

Press Exit when finished with the Settings screen to continue to the Confirm screen.

If the displayed settings are as desired, press OK to continue to the Clinician Home Screen. Otherwise, press Back to change any incorrect settings.

If settings were changed with V.A.C.* Therapy off, press the Therapy On/Off button on the Clinician Home Screen to start therapy.
The Settings Guide helps the clinician select from pre-set therapy ranges according to wound type and treating physician’s orders. Selected ranges are a guide based on common settings for different wound types. Individual patient conditions may vary. Consult physician to verify settings for each patient.

Should physician orders fall outside the pre-set therapy ranges, select Other in this mode or use Manual Therapy Settings detailed in the Settings Screen (Manual) section of this manual.

Select Wound Type Screen

From the Clinician Home Screen press Therapy, then OK, then Settings Guide, then OK to access the Select Wound Type screen.

Use the + and - buttons to scroll through the available wound type selections.

Press Cancel on any screen to cancel and exit the procedure.

Select Pressure Screen

When finished with the Select Wound Type screen, press Next to continue to the Select Pressure screen.

Use the + and - buttons to scroll through the pressure selections.

Pressure selections are in ranges for the wound type selected on the previous screen.
Select Mode Screen

When finished with the Select Pressure screen, press Next and move to the Select Mode screen.

Use the + and - buttons to choose Continuous or Intermittent Therapy.

When finished with this screen, press Next.

If Intermittent is not an option for the wound type selected, the mode will not change.

Intermittent can be manually set from the Settings screen, or press Back and select Other as the wound type.

Intermittent Type Screen

If Intermittent Therapy was chosen on the previous screen, this screen will appear.

Use the + and - buttons to change the desired On and Off Time (in minutes).

Both On and Off Times can be set from 1 minute to 10 minutes in 1 minute increments.

When finished with this screen, press Next.

Default is: On Time = 5 minutes, Off Time = 2 minutes.

Settings Guide Confirmation

Once the settings are chosen, the Confirm screen will appear.

If the displayed settings are as desired, press OK to continue to the Therapy Screen. Otherwise, press Back to change any incorrect settings.

Settings take effect when OK is pressed.

Settings Guide Intensity default is Low. Intensity can only be changed using the Manual Therapy Settings (see Settings Screen (Manual) section).
SEAL CHECK™ Screen

The **SEAL CHECK™** screen provides a visual and audible representation of the air flow rate in the SENSAT.R.A.C.™ system. For more information about the SEAL CHECK™ Feature see **SEAL CHECK™ Feature Overview** section.

From the **Clinician Home Screen** press **Therapy**, then **OK**, then **SEAL CHECK™ Button** to access the **SEAL CHECK™ Feature screen**.

Press **Seal Audio** to turn the Seal Audio tone on or off.

This tool can be used as an aid to ensure that the dressing system is correctly sealed.

Therapy History Screen

The **Therapy History** screen displays the therapy information in date, time and event columns. The date is in ascending order and time is displayed using the 24-hour clock format.

From the **Clinician Home Screen** press **Therapy**, then **OK**, then **Therapy History** to access the **Therapy History** screen.

Use the + and - buttons to scroll to additional listings.

If wound history is recorded, see **Patient History** screen for wound progress history.

Press to return to the **Therapy** screen.

Patient History Screens

The Patient History screens enable a clinician to:

- Create a new access code.
- Delete the current patient record.
- Start a new patient record.
- View the patient record.
- Export history.
- View a wound image area graph.
- Analyze wound images.
Enter Access Code Screen

From the Clinician Home Screen press Therapy, then OK, then Patient History to access the Enter Access Code screen.

Press to proceed to the next screen.

Press to create a new access code.

Press to return to the Therapy screen.

This icon appears in the Therapy Status Bar and Display Area when a Patient Record is active.

For security purposes:

- The INFOV.A.C.™ Therapy Unit will only allow one patient record at a time to be active. If a new access code is entered, the current access code is overwritten and all patient history associated with it is deleted.

  All information will be automatically deleted when the unit is returned to KCI.

- If a current access code is incorrectly entered, access to the Patient History screen will be denied and you will automatically be returned to the Therapy screen.

- If an incorrect access code is entered ten times, the system will lock. Contact your local KCI representative.

To create a new access code:

1. Press Create.

2. Enter your four digit numerical access code using the numbers on the screen, then press OK.

3. Re-enter your four digit numerical access code.

4. Press OK to set the new access code and proceed to the Change Patient ID screen.
Change Patient ID Screen

A unique number is automatically generated from the date and time unless a custom ID is created.

1. Enter the new Patient ID on keyboard if desired.

2. Press OK when you have entered the new Patient ID, or are ready to proceed to the Patient History screen.

3. Press Exit to continue to the Confirm screen.

Press this button as a space bar.

Press this button to backspace/delete.

If no changes are made, the automatically generated number will be used as the Patient ID.

Patient History Screen

From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK to access the Patient History screen.

Press to view Patient History and add notes (up to 30 characters).

Press to delete a Patient History file.

Press to export Patient History and Therapy History.

Press to view a graphical representation of measured wound area over time.

Press to adjust Leak Alarm settings.

Press to view and perform Wound Image Analysis.
View History Screen

From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then View History to access the Patient History screen.

The Patient History screen will display short notes (up to 30 characters) added by the clinician and/or medical staff. For a new Patient History this screen will be blank.

Add Note To History Screen

From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then View History then Add Note on the Patient History screen to open the Add Note to History screen.

Use the screen keys to enter a note (up to 30 characters). The notes entered will be displayed in the window area above the keys.

Press to log new note and return to Patient History screen.

Press this button as a space bar.

Press this button to backspace/delete.

Press to return to the Patient History screen.

Delete History Screen

From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then Delete History to access the Delete History screen.

Deleted History cannot be recovered.

Press and hold Delete for two seconds to delete the Patient History Records. After the Patient History Records have been deleted, the screen will return to the Therapy screen.

Press to return to the Patient History screen without deleting Patient History.

All information will be automatically deleted when the unit is returned to KCI.
Export Patient History Screen

If Unit has an IR Port

From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then Export History to access the Export Patient History screen.

The exported Patient History includes folders containing Therapy History, Patient History and all wound images for the current patient.

Image files are exported with the file names created during the imaging process.

Files can be exported to a Memory Card, USB device or IR Port (if unit is equipped).

Access the Memory Card Transfer screen.
Access the USB Transfer screen.
Access the IR Transfer screen.

If Unit Does Not Have an IR Port

The connection ports are for clinician use only.
The Memory Card Slot is for authorized clinician access only.

1. Insert the Memory Card into the INFOV.A.C.™ Therapy Unit Memory Card Slot. An audible click signals that the card is properly inserted.

2. From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then Export History, then Export to Memory Card to access the Memory Card Transfer screen.

3. Press Next to continue.

A progress bar on the screen will indicate transfer status.

4. When the transfer is complete press OK to continue.

If the Memory Card Transfer Error screen appears when the Memory Card is inserted into the Memory Card Slot, files could not be transferred.

Possible reasons for transfer error:

- Card not inserted correctly.
- Incorrect format card.
- No images in the system.
The USB port is for authorized clinician access only. USB devices should be connected directly and only unpowered USB mass storage devices should be attached to the INFOV.A.C.™ Therapy Unit. AC or battery powered drives, computers, computer equipment, other devices or USB extension leads should not be connected to this device.

1. Insert the USB device into the INFOV.A.C.™ Therapy Unit USB port.

2. From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then Export History, then Export to USB to access the USB Transfer screen.

3. Press Next to continue.

A progress bar on the screen will indicates transfer status.

4. When the transfer is complete press OK to continue.

If the USB Transfer Error screen appears when the USB device is inserted into the USB port, files could not be transferred.

Possible reasons for transfer error:
- The USB device is not inserted correctly.
- The incorrect type of device is connected.
IR Transfer Screen

IR Transfer

Open data door on InfoV.A.C.® therapy unit, place IR device in front of IR window and select ‘Next’

1. From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then Export History, then Export to IR to access the IR Transfer screen.

2. Open the Data Door and position the IR device in front of the INFOV.A.C.™ Therapy Unit.

3. Aim the IR Data Port of the transfer device at the Infrared Data Port on the INFOV.A.C.™ Therapy Unit.

4. Press Next to continue.

The INFOV.A.C.™ Therapy Unit will attempt to establish a connection with the transfer device; once the connection is established, the transfer will begin.

A progress bar on the screen will indicates transfer status.

5. When the transfer is complete press OK to continue.

If the IR Transfer Error screen appears, the INFOV.A.C.™ Therapy Unit does not detect the IR device or the device is incompatible. Files could not be transferred.

Possible reasons for transfer error:

- The IR transfer device is not lined up with INFOV.A.C.™ Therapy Unit’s Infrared Data Port.
- The IR transfer device’s IR port is not active or is not set to receiving mode.
- Lenses on the IR device are damaged or obscured.
**Graph Screen**

From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then View Graph to access the Graph screen.

The graph displays a representation of image area over time.

A graph cannot be constructed if the Patient History file has been deleted.

A graph cannot be constructed unless measurements of the image area have been previously saved in the patient's history log. At least two measurements from different days are required (area of the image against time) for a graph to be constructed.

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**Select Leak Alarm Threshold Screen**

Consult the clinician prescribing therapy before making changes to this setting.

From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then Alarm Adjust to access the Select Leak Alarm Threshold screen.

Use the + and - buttons to select the level of alarm adjustment required.

The Leak Alarm Threshold is set by default to High (2 Liters per minute), this screen allows a change to Low (1 Liter per minute) when appropriate.

Press to confirm the alarm adjustment settings.
Wound Image Analysis

The Wound Imaging feature is an aid in recording treatment progress.

**Wound imaging area and volume calculation features are not intended to be exact measurements and are not intended for use in the diagnosis and treatment of wounds.**

Accessories required to utilize this feature include:

- Digital camera that uses an SD/MMC card. The maximum size of the SD/MMC card that can be used to upload images to an INFOV.A.C.™ Therapy Unit is 2 GB. The unit will not upload images using a SD/MMC card larger than 2 GB. Please check the size of the SD/MMC card prior to uploading images into the INFOV.A.C.™ Therapy Unit.

  The maximum picture resolution accepted by the INFOV.A.C.™ Therapy Unit is 2.2 megapixels. If using a camera with higher resolution, you must adjust the picture resolution settings on your camera to a setting of 2.2 megapixels or less. Please refer to your camera user manual for further instructions.

- An SD/MMC card.

- Calibration Reference Square. This is found on the ruler in the V.A.C.® Dressing package. This reference square is needed for the INFOV.A.C.™ Therapy Unit to calculate wound measurements.

- Stylus. This is found inside the data door (see INFOV.A.C.™ Therapy Unit - Parts Identification section). This is needed for tracing.

For optimal operation of the Wound Image Analysis feature it is recommended that:

- The sterile Calibration Reference Square should be placed in the same location on the wound each time an image is taken.

- All images be taken from directly above the wound.

- The wound and Calibration Reference Square should fill as much of the image as possible.

- The image should be taken in good light conditions.

- Images must be taken directly from the camera, not transferred or downloaded to the card from a computer.

**Using a camera that has a date and time function will allow for easier tracking of images.**
Troubleshooting Suggestions:

- **Media Image Capacity:** Keep as few images on the media as possible prior to uploading the images into the INFOV.A.C.™ Therapy Unit. It is recommended you upload any existing pictures from your media (SD/MMC card or USB memory stick) to your computer prior to capturing new wound photos on the media. By having too many images on the media, you run the risk of receiving a “DSP Not Initialized” error which will freeze the unit, preventing image upload.

- **“DSP Not Initialized” Error:** This indicates an issue with the image uploading and prevents the unit from uploading images into the INFOV.A.C.™ Therapy Unit. If you receive this error, exit to the previous screen and reset the unit by turning the unit off and back on again. If you cannot go back to the previous screen, reset the unit by turning the unit off and back on again. You may then proceed to the image uploading screen as directed in this manual (see *Imaging Screen* section).

- **Screen Freezes during Image Uploading:** This indicates an issue with the image uploading into the INFOV.A.C.™ Therapy Unit. If you receive this error, reset the unit by turning the unit off and back on again. You may then proceed to the image uploading screen as directed in this manual (see *Imaging Screen* section).

- **Erasing Images from the Unit:** Digital images cannot be individually erased from the INFOV.A.C.™ Therapy Unit. When the internal memory is full and more pictures need to be uploaded, the patient history must be cleared by deleting the record and creating a new patient history. It is recommended that the patient history record be exported prior to deleting so all history and images are saved. See *Patient History Screens* section of this manual for further instructions on viewing, deleting, exporting, and creating a patient history record.
Imaging Screen

From the Clinician Home Screen press Therapy, then OK, then Patient History, then enter or create an access code and press OK, then Imaging to access the Imaging screen.

Press to select and view an image from the SD/MMC card.

Press to select and view an image from a USB device.

Press to select and view an image from the INFOV.A.C.™ Therapy Unit internal memory.

There will be a short delay while the images are accessed from the internal memory or the memory card. The relevant button will be highlighted with a green rim to indicate that this is in progress.

When the INFOV.A.C.™ Therapy Unit is ready, the File Select screen will appear.

If INFOV.A.C.™ Memory is selected for a new Patient Record, no images will be available and the image thumbnail will be black.
File Select Screen

The number of files available for viewing and the number of the currently-viewed file (for example, 9 of 10) is displayed in the upper right of the File Select screen.

Use the + and - buttons to select the next or the previous image.

Press Select when the required image is displayed.

Images being opened from a memory card or USB device will be automatically downloaded into the INFOV.A.C.™ Therapy Unit internal memory.

Press to begin the Wound Area Trace procedure.

Press to calculate the wound area inside the highlighted trace outline.

Press to begin the Wound Volume calculation procedure.
Image Reference Calibration

Trace the reference square in the image to scale the image area and volume measurements.

- On-screen instructions are provided at the bottom of each screen to guide the user through the process.
- Use only the stylus provided to trace the reference square.
- Press to begin the Calibration Reference process.

Touch each corner of the reference square. When the last corner of the reference square is touched the corner points will be joined by a highlighted line.

1. 2. 3. 4.

It is important to select corners in either a clockwise or counter-clockwise manner. Incorrect sequence will lead to a calibration error.

If necessary, press to restart the Calibration Reference process.

Wound Area Trace

Next, trace the outline of the wound area. Press the button pictured at left to begin the Wound Area Trace procedure. The Select Image Area screen will appear.

Press Image Area 1 to trace the overall outline of the wound area. All traces and results for Image Area 1 will be displayed in yellow.

The INFOV.A.C.™ Therapy Unit only stores the area and depth of Image Area 1 in the Patient Log. This information in the Patient Log is used to construct a graph.

Press Image Area 2 to trace another area inside the wound. All traces and results for Image Area 2 will be displayed in blue.

Press Image Area 3 to trace an additional area. All traces and results for Image Area 3 will be displayed in pink.
Once an Image Area is selected, the image will be displayed as a full screen.

Draw around the required area with the stylus.

- A yellow square will appear at the start point. The trace is completed when the end of the yellow line returns to the start point.

To correct an error, tap the stylus in the square at the start point. This action will return you to the previous screen where you can select to retrace the area.

   When the outline of the wound image area is complete, the buttons will reappear.

   If required, press Wound Trace Area button again to trace additional areas.

Calculate Wound Area

Press the button pictured at left to calculate the wound area inside the highlighted trace outline.

The results will be displayed in the color of the trace for the appropriate measured area.
Calculate Wound Volume

Select Image Area for Depth Screen

The wound volume can be calculated from the areas previously traced and the measured depth.

Press the button pictured at left to begin the Wound Volume calculation procedure. The Select Image Area for Depth screen will appear.

The INFOV.A.C.™ Therapy Unit only stores the area and depth of Image Area 1 in the Patient Log. This information in the Patient Log is used to construct a graph.

Press to enter the depth of Image Area 1.

Press to enter the depth of Image Area 2.

Press to enter the depth of Image Area 3.

Use the screen keys to enter the image depth.

Press OK when finished entering image depth.

Image area and volume will be displayed on the screen at left.

Press Exit to return to the Patient History screen.
Clinician Utilities

From the Clinician Home Screen, press Utilities to access the Utilities screen.

From this Utilities screen the clinician can:
- Set the time and date to current time and calendar date.
- Adjust regional settings.
- Set the brightness of the touch screen user interface. The green crescent at the bottom of the button indicates the level of brightness:
  - Low
  - Medium
  - High

The Default setting is Medium

Press to return to the Clinician Home Screen.

Changing Time and Date

From the Clinician Home Screen press Utilities, then press Time/Date to access the Set Time/Date screen.

Press to set current local time and calendar date. Holding these buttons will rapidly scroll through available selections.

Press to return to the Utilities screen.
Changing Pressure Units and Date Format

From the Clinician Home Screen press Utilities, then press Regional Settings to access this screen.

Press to switch between mmHg (millimeters of mercury) and kPa (kilo-pascals) units of measurement.

The INFOV.A.C.™ Therapy Unit is designed to show two units of measure with mmHg (millimeters of mercury) being the default. If you prefer kPa (kilo-Pascals), press Pressure Units to switch from mmHg to kPa.

Press to change how decimals are displayed.

Press to select cm (centimeters) or in (inches)

Press to change the language in which the screens are displayed.

Press to switch between displaying DD/MMM/YYYY (Day-Month-Year) and MMM/DD/YYYY (Month-Day-Year) formats.

Default settings are: mmHg, DD/MMM/YYYY, cm and English.

Press Exit to return to the Utilities screen.
Alerts and Alarms

ATTENTION: Important Information about Alerts and Alarms

A **Low Priority Alarm / Alert** will be displayed on the touch screen when the INFOV.A.C.™ Therapy Unit detects a condition that requires patient or caregiver attention.

Alerts will be accompanied by a **single** audible tone.

A **Medium Priority Alarm** will be displayed on the touch screen user interface when the INFOV.A.C.™ Therapy Unit detects a condition that **requires immediate patient or caregiver attention in order to ensure the prescribed therapy is being delivered**.

Alarms will be accompanied by a **repeating** audible tone.

If alarm conditions cannot be resolved, contact KCI.

Press **Audio Pause** to silence the audible tone for two minutes.

Press **Help** for more information about the alert or alarm.

**WARNING:** According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.

This applies to all alarm conditions when therapy is interrupted.
Battery Low Alert
Low Priority Alert - This alert screen indicates approximately two hours of battery power remain. This alert will be accompanied by a single audible tone.

This alarm may be silenced during troubleshooting by pressing Audio Pause.

To resolve this alert:

1. Connect therapy unit to wall outlet using the INFOV.A.C.™ Power Supply to recharge battery. An amber light above the power connection will indicate unit is charging. Refer to the Charging the Battery section of this manual for more information.

2. Press Reset on this screen to return to the Home Screen.

Battery Critical Alarm
Medium Priority Alarm - This alarm screen indicates approximately 30 minutes of battery power remain. This alarm will be accompanied by a repeating audible tone.

This alarm may be silenced during troubleshooting by pressing Audio Pause.

To resolve this alarm:

1. Connect therapy unit to wall outlet using the INFOV.A.C.™ Power Supply to recharge battery. An amber light above the power connection will indicate unit is charging. Refer to the Charging the Battery section of this manual for more information.

2. Press Reset on this screen to return to the Home Screen.

3. Ensure therapy is on by confirming that the green crescent is lit on the Therapy On/Off button. If not, press the Therapy On/Off button to restart therapy.

V.A.C.® Therapy continues; however, if this alarm is not resolved within one minute, therapy will be interrupted.
Canister Full Therapy Interrupted Alarm

Medium Priority Alarm - This alarm screen appears when the canister is full and should be replaced. This alarm will be accompanied by a repeating audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing Audio Pause.

To resolve this alarm:

1. Determine if canister is full by comparing the level of fluid to the graduated marks on the canister (see below).

2. If canister is not full, press Reset.

3. If canister is full, change canister and press Reset to return to the Home Screen.

4. Restart therapy by pressing Therapy On/Off on the Home Screen.

See the Changing the Canister section of this manual for additional information.

WARNING: According to clinician instructions, replace V.A.C.* Dressing with alternate dressing if therapy is off for more than two hours.
Canister Not Engaged Alarm

Medium Priority Alarm - This alarm screen appears when the canister is not fully inserted and/or properly latched. This alarm will be accompanied by a repeating audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing Audio Pause.

To resolve this alarm:

1. Remove the canister by pressing the Canister Release button.

2. Inspect the canister and INFOV.A.C.™ Therapy Unit to ensure that foreign objects or debris do not interfere with the canister and therapy unit’s mating surfaces.

3. Ensure both seals are present. If seals are missing or damaged, contact KCI.

4. Re-attach the canister to the INFOV.A.C.™ Therapy Unit ensuring that the canister is fully engaged and latched. An audible click indicates that the canister is properly installed.

5. Press Reset on this screen to return to the Home Screen.

6. Restart therapy by pressing Therapy On/Off.

7. If this alarm continues to appear, repeat steps 1 through 6 with a new canister.

If alarm condition cannot be resolved, contact KCI.

WARNING: According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.
Leak Alarm

Medium Priority Alarm - This alarm screen appears when a significant negative pressure leak has been detected. If this alarm is not resolved in three minutes, therapy will be interrupted. This alarm will be accompanied by a single audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing Audio Pause.

To resolve this alarm:

1. Ensure connector between dressing tubing and canister tubing is properly locked.

2. Ensure canister is fully engaged. (See Canister Not Engaged Alarm section).

3. Press the SEAL CHECK™ Button to use the SEAL CHECK™ Feature to help identify leaks in dressing. Refer to the Finding a Leak Using SEAL CHECK™ section of this manual for details on how to use the SEAL CHECK™ Feature and how to repair leaks.

4. Refer to the Dressing Application Instructions for Use provided with V.A.C.* Dressings for information on using excess V.A.C.* Drape material to seal the leak area.

5. Once the leak has been resolved, press Reset on this screen to return to the Home Screen.

6. Ensure therapy is on by confirming that the green crescent is lit on the Therapy On/Off button. If not, press the Therapy On/Off button to restart therapy.

If this alarm is not resolved within three minutes, the Leak Alarm Therapy Interrupted alarm will appear and therapy will stop.

Refer to Leak Alarm Therapy Interrupted section for procedures to restart therapy.

WARNING: According to clinician instructions, replace V.A.C.* Dressing with alternate dressing if therapy is interrupted or off for more than two hours.
Leak Alarm Therapy Interrupted

Medium Priority Alarm - This alarm screen appears when a detected negative pressure leak has not been resolved and therapy has been interrupted. This alarm will be accompanied by a repeating audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing Audio Pause.

To resolve this alarm:

1. Press Reset on this screen to return to the Home Screen.
2. Restart therapy by pressing Therapy On/Off.

If the leak condition is not resolved, access the SEAL CHECK™ tool for further troubleshooting:

- From the Clinician Mode Screen, Therapy On/Off opens the SEAL CHECK™ screen on the Therapy Start screen.
- Or
- From the Clinician Home Screen press Therapy, then OK, then SEAL CHECK™ to access the SEAL CHECK™ Feature.

The patient’s only access to the SEAL CHECK™ Feature is only through the Leak Alarm screen.

If the leak condition is not resolved within three minutes, the alarm screen will once again appear and therapy will stop.

If alarm condition cannot be resolved, contact KCI.

WARNING: According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.
Blockage Alert

Low Priority Alarm - This alert screen appears when the INFOV.A.C.™ Therapy Unit has detected a potential blockage. This alert will be accompanied by a single audible tone.

To resolve this alert:

1. Ensure both clamps on the dressing and canister tubing are open.

2. Ensure that the tubing is not kinked, crimped or blocked in any way.

3. If the Blockage Alert remains after completing steps 1 and 2, lower the therapy unit and tubing to be level with or below the wound site. If the alert is resolved by lowering the unit, normal use may resume.

4. Press Reset on this screen to return to the Home Screen.

The INFOV.A.C.™ Therapy Unit will continue to attempt to apply therapy during this alert.
Blockage Alarm Therapy Interrupted

Medium Priority Alarm - This alarm screen appears when the INFOV.A.C.™ Therapy Unit has determined that a blockage is present. This alarm will be accompanied by a **repeating** audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing **Audio Pause**.

To resolve this alarm:

1. Ensure both clamps on the dressing and canister tubing are open.

2. Ensure that the tubing is not kinked, crimped or blocked in any way.

3. If the Blockage Alarm Therapy Interrupted remains after completing steps 1 and 2, lower the therapy unit and tubing to be level with or below the wound site. If the alert is resolved by lowering the unit, normal use may resume.

4. Press **Reset** on this screen to return to the **Home Screen**.

**WARNING:** Therapy unit remains on; however, negative pressure at the wound is below therapeutic value.

**WARNING:** According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.

If alarm condition cannot be resolved, contact KCI.
Low Pressure Alert

Low Priority Alert - This alert screen appears when the INFOV.A.C.™ Therapy Unit has not reached the selected therapy set pressure. This alert will be accompanied by a single audible tone.

To resolve this alert:

1. Ensure both clamps on the dressing and canister tubing are open.

2. Ensure that the tubing is not kinked, crimped or blocked in any way.

3. If the Low Pressure Alert remains after completing steps 1 and 2, lower the therapy unit and tubing to be level with or below the wound site. If the alert is resolved by lowering the unit, normal use may resume.

4. Press Reset to return to the Home Screen.

V.A.C.® Therapy is still being applied, but at a lower than selected pressure.
Low Pressure Alarm Therapy Interrupted

Medium Priority Alarm - This alarm screen appears when the INFOV.A.C.™ Therapy Unit has not reached the selected therapy set pressure and negative pressure at the wound is below therapeutic value. This alarm will be accompanied by a repeating audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing Audio Pause.

To resolve this alarm:

1. Ensure both clamps on the dressing and canister tubing are open.

2. Ensure that the tubing is not kinked, crimped or blocked in any way.

3. If the Low Pressure Alarm Therapy Interrupted remains after completing steps 1 and 2, lowering the therapy unit and tubing to be level with or below the wound site may resolve this alarm. If the alert is cleared by lowering the unit, normal use may resume.

4. Press Reset on this screen to return to the Home Screen.

WARNING: Therapy unit remains on; however, negative pressure at the wound may be below set pressure, potentially compromising therapeutic benefits.

WARNING: According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.

If alarm condition cannot be resolved, contact KCI.
Therapy Inactive Alarm

Medium Priority Alarm - This alarm screen appears when V.A.C.® Therapy has been off for fifteen minutes (with the unit powered on) without the touch screen being touched. This alarm will be accompanied by a repeating audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing Audio Pause.

To resolve this alarm:

1. Press Reset on this screen to return to the Home Screen.
2. Restart therapy by pressing Therapy On/Off.
3. If V.A.C.® Therapy is not desired, turn the INFOV.A.C.™ Therapy Unit off by using the Power On/Off button on the front of the unit.

WARNING: According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.

System Error Alarm Therapy Interrupted

Medium Priority Alarm - This alarm screen appears when there is a technical fault within the INFOV.A.C.™ Therapy Unit. Several different types of system errors may occur. A number will appear in the yellow alarm box that represents the diagnostic code of the technical fault. This alarm will be accompanied by a repeating audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing Audio Pause.

To resolve this alarm:

1. Record the error number.
2. Power the unit off and then on using the Power On/Off button on the front of the unit.
3. If error persists, contact KCI.

WARNING: According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.
Service Timer Expired Alert

Low Priority Alert - This alert screen appears when the INFOV.A.C.™ Therapy Unit has reached its service time limit. Once the Service Timer has expired, this alert will appear every time the unit is powered on. When Days Left reaches zero, this alert will reappear periodically during therapy.

To resolve this alert:
1. Contact your local KCI representative to obtain a new service timer code.
2. Press Enter Code to enter the code obtained from KCI.

Internal Temperature Alarm

Medium Priority Alarm - This alarm screen appears when the internal temperature of the INFOV.A.C.™ Therapy Unit has reached its maximum limit. This alarm will be accompanied by a repeating audible tone.

To resolve this alarm:
1. Move the INFOV.A.C.™ Therapy Unit to a cooler location.
2. Press Reset on this screen to return to the Home Screen.

Therapy continues while this alarm is active. The screen will be turned off after five minutes, but will illuminate when touched. Battery charging is stopped.

If alarm condition cannot be resolved, contact KCI.

WARNING: According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.
Pressure Deviation Therapy Interrupted

Medium Priority Alarm - This alarm screen appears when the wound pressure has deviated above the system limits. This alarm will be accompanied by a repeating audible tone.

This alarm may be silenced for two minutes during troubleshooting by pressing Audio Pause.

To resolve this alarm:

Power the unit off and then on using the Power On/Off button on the front of the unit.

If alarm condition cannot be resolved, contact KCI.

WARNING: According to clinician instructions, replace V.A.C.® Dressing with alternate dressing if therapy is interrupted or off for more than two hours.
Care and Cleaning

Standard Precautions
The following are the KCI recommended daily and weekly cleaning and infection control procedures for the INFOV.A.C.™ Therapy Unit.

![Always follow Standard Precautions.](image)

Standard Precautions are designed to reduce the risk of transmission of microorganisms from both known and unknown sources of infection. These precautions can be applied to all patients, regardless of their diagnosis or presumed infection status, and should be used when contact is anticipated with blood and all body fluids. This also includes secretions and excretions (except sweat) regardless of whether blood is visible or not, non-intact skin (i.e., open wounds) and mucous membranes.

Waste Disposal
Discard all disposable items (all tubing, connectors, clamps, used canister, used dressings, etc.) in accordance with local medical waste disposal regulations. Improper disposal may run the risk of regulatory non-compliance.

Cleaning the INFOV.A.C.™ Therapy Unit
Cleaning and disinfection of the INFOV.A.C.™ Therapy Unit includes wipedown of all hard surface components. Follow your institutional procedures used for cleaning and disinfection of other hard surface durable electronic medical equipment. The INFOV.A.C.™ Therapy Unit must be cleaned and disinfected:

- If it becomes soiled during patient use.
- At least weekly.
- In between patients.

![Ensure that the INFOV.A.C.™ Therapy Unit is powered off and disconnected from AC power when using cleaning fluids of any nature.](image)

KCI recommends the following regarding cleaning and disinfecting KCI V.A.C.* Therapy devices:

- To help reduce risk of infection and contact with blood and body fluids, use personal protective equipment (PPE) such as medical procedure gloves.
- Clean all organic material (visible soil or body secretions) from the therapy unit prior to disinfection.
- Use hospital-grade cleaners and disinfectants.
• Do not immerse or saturate the therapy unit with fluids to avoid damage to the electronics in the device.

• Do not use alcohol based solutions around the touchscreen edges or near gasket and power switches since alcohol based solutions will easily wick up into the screen and may cause equipment malfunction.

Cleaning the Touch Screen

1. Press the Screen Guard button on the Home screen (refer to Patient Mode Control Screens section) to activate Screen Guard. The lock button icon will close and the Screen Guard screen will appear.

2. Use a soft, non-abrasive cloth to gently clean the touch screen.

   Do not use any liquid to clean the touch screen.

   Do not use excessive force to clean the touch screen. Pressing too hard may cause damage.

3. To unlock the touch screen, press the 1 button, then the 2 button on the Screen Guard screen to return to the Home screen.
Electromagnetic Compatibility (EMC)

The following are guidance and manufacturer’s declarations regarding EMC for the INFOV.A.C.™ Therapy Unit.

- The INFOV.A.C.™ Therapy Unit needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in the following pages.

  *This equipment is intended for use by healthcare professionals only. As with all electrical medical equipment, this equipment 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 INFOV.A.C.™ Therapy Unit or shielding the location.*

- The emissions characteristics of this equipment make it suitable for use in industrial areas and hospitals (CISPR 11 class A). If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radio-frequency communication services.

- Portable and Mobile RF communications equipment, RFID readers, electronic article surveillance (anti-theft) equipment and metal detectors can affect the performance of the INFOV.A.C.™ Therapy Unit. Please use the guidelines and recommendations specified in Tables 204 and 206.

- Other medical equipment or systems can produce electromagnetic emissions and therefore can interfere with the functionality of the INFOV.A.C.™ Therapy Unit. Care should be used when operating the INFOV.A.C.™ Therapy Unit adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, the INFOV.A.C.™ Therapy Unit and the other equipment should initially be observed to verify normal operation in the configuration in which it will be used.

- The electrical cables, external power supplies and accessories listed or referenced in this manual have been shown to comply with the test requirements listed in the following tables. Care should be taken to use only manufacturer-recommended cables, power supplies and accessories with the INFOV.A.C.™ Therapy Unit. If a third-party supplier offers cables, external power supplies and electrical accessories for use with the INFOV.A.C.™ Therapy Unit and they are not listed or referenced in this manual, it is the responsibility of that third-party supplier to determine compliance with the standards and tests in the following tables.

- The use of electrical cables and accessories other than those specified in this manual or referenced documents may result in increased electromagnetic emissions from the INFOV.A.C.™ Therapy Unit or decreased electromagnetic immunity of the INFOV.A.C.™ Therapy Unit.
Table 201

Guidance and Manufacturer’s Declaration – Electromagnetic Emissions

The INFOV.A.C.™ Therapy Unit is intended for use in the electromagnetic environment specified below. The customer or the end user of the INFOV.A.C.™ Therapy Unit should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions test</th>
<th>Compliance</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF Emissions - CISPR 11 (Radiated &amp; Conducted)</td>
<td>Group 1</td>
<td>The INFOV.A.C.™ Therapy Unit uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>RF Emissions - CISPR 11 (Radiated &amp; Conducted)</td>
<td>Class A</td>
<td>The INFOV.A.C.™ Therapy Unit is suitable for use in all establishments, other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonic Emissions IEC 61000-3-2</td>
<td>Class A</td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations / Flicker Emissions IEC 61000-3-3</td>
<td>Complies</td>
<td></td>
</tr>
</tbody>
</table>

Table 202

Guidance and Manufacturer’s Declaration – Electromagnetic Immunity

The INFOV.A.C.™ Therapy Unit is intended for use in the electromagnetic environment specified below. The customer or the end user of the unit should assure it is used only in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>EN / IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Intended Electromagnetic Environment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic Discharge (ESD) IEC 61000-4-2</td>
<td>± 6kV contact</td>
<td>± 8kV contact</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td></td>
<td>± 8kV air</td>
<td>± 15kV air</td>
<td></td>
</tr>
<tr>
<td>Electric fast transient / burst IEC 61000-4-4</td>
<td>± 2kV for power supply lines</td>
<td>± 2kV for power supply lines 100 kHz repetition frequency</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td></td>
<td>± 1kV for input / output lines</td>
<td>± 1kV for input / output lines 100 kHz repetition frequency</td>
<td></td>
</tr>
<tr>
<td>Surge IEC 61000-4-5</td>
<td>± 1kV differential mode (line-line)</td>
<td>± 1kV differential mode (line-line)</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td></td>
<td>± 2kV common mode (line-earth)</td>
<td>± 2kV common mode (line-earth)</td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11</td>
<td>&lt;5% Ut (&gt;95% dip in Ut) for 0.5 cycle</td>
<td>Dips: 0% Ut for 1 cycle 70% Ut for 25 cycles at 50Hz or 30 cycles at 60Hz</td>
<td>Product has internal battery backup. If the user of the INFOV.A.C.™ Therapy Unit requires continued operation during power mains interruptions, it is recommended that the unit be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td></td>
<td>40% Ut (60% dip in Ut) for 5 cycles</td>
<td>Single phase: at 0°</td>
<td></td>
</tr>
<tr>
<td></td>
<td>70% Ut (30% dip in Ut) for 25 cycles</td>
<td>Interruptions: 0% Ut for 250 cycles at 50Hz or 300 cycles at 60Hz</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt;5% Ut (&gt;95% dip in Ut) for 5 seconds</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Power frequency (50/60Hz) magnetic field IEC 61000-4-8</td>
<td>3 A / m</td>
<td>30 A / m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

NOTE: Ut is the AC mains voltage prior to application of the test level.
**Table 204**

**Guidance and Manufacturer’s Declaration – Electromagnetic Immunity**

The INFOV.A.C.™ Therapy Unit is intended for use in the electromagnetic environment specified below. The customer or the end user of the unit should assure it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>EN / IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td></td>
<td></td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the INFOV.A.C.™ Therapy Unit, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
<tr>
<td>IEC 61000-4-6</td>
<td>3Vrms 150kHz to 80MHz</td>
<td>3Vrms 150kHz to 80MHz</td>
<td>d = 1.2√P 80MHz to 800 MHz</td>
</tr>
<tr>
<td>Radiated RF</td>
<td>3V / m 80MHz to 2.5GHz</td>
<td>6Vrms in ISM bands between 150kHz to 80MHz 80% AM at 1kHz 3V / m 80MHz to 2.7GHz</td>
<td>d = 2.3√P 800MHz to 2.5GHz where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended minimum separation distance in meters (m).</td>
</tr>
</tbody>
</table>

Field strengths from fixed RF transmitters, as determined by an electromagnetic site surveya, should be less than the compliance level in each frequency rangeb. Interference may occur in the vicinity of equipment marked with the following symbol:

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

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

Field strengths from fixed transmitters, such as base stations for radio (cellular / cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the INFOV.A.C.™ Therapy Unit is used exceeds the applicable RF compliance level above, the unit should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the INFOV.A.C.™ Therapy Unit.

NOTE: This equipment has been tested and found to comply with the limits for medical devices to IEC 60601-1-2: 2014 4th edition. These limits and test levels are intended to provide reasonable safety with regard to electromagnetic disturbances when the device is used in a typical medical installation.

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*a Field strengths from fixed transmitters, such as base stations for radio (cellular / cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the INFOV.A.C.™ Therapy Unit is used exceeds the applicable RF compliance level above, the unit should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the INFOV.A.C.™ Therapy Unit.

*b Over the frequency range 150kHz to 80MHz, field strengths should be less than 3V / m.
Table 206

Recommended separation distances between portable and mobile RF communications equipment and the INFOV.A.C.™ Therapy Unit

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

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter in watts (W)</th>
<th>Separation distance according to frequency of transmitter in meters (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150kHz to 800MHz</td>
</tr>
<tr>
<td></td>
<td>80MHz to 800MHz</td>
</tr>
<tr>
<td></td>
<td>800MHz to 2.5GHz</td>
</tr>
<tr>
<td></td>
<td>(d = 1.2\sqrt{P})</td>
</tr>
<tr>
<td>0.01</td>
<td>.12</td>
</tr>
<tr>
<td>0.1</td>
<td>.38</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.8</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
</tbody>
</table>

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

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

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

Included Power Supplies

<table>
<thead>
<tr>
<th>Part Number</th>
<th>Description</th>
<th>Manufacturer</th>
<th>Max Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>M6266106</td>
<td>Power Supply</td>
<td>Astec Power or ICC</td>
<td>3.05 m</td>
</tr>
<tr>
<td>M4268882</td>
<td>Elec / Con-US Cordset - INFOV.A.C.™</td>
<td>Electri-Cord Mfg. Co.</td>
<td>2.03 m</td>
</tr>
<tr>
<td>M4268883</td>
<td>Elec / Con-Australia Cordset - INFOV.A.C.™</td>
<td>Electri-Cord Mfg. Co.</td>
<td>2.05 m</td>
</tr>
<tr>
<td>M4268880</td>
<td>Elec / Con-UK Cordset - INFOV.A.C.™</td>
<td>Electri-Cord Mfg. Co.</td>
<td>2.05 m</td>
</tr>
<tr>
<td>M4268881</td>
<td>Elec / Con-EU Cordset - INFOV.A.C.™</td>
<td>Electri-Cord Mfg. Co.</td>
<td>2.05 m</td>
</tr>
<tr>
<td>4103864</td>
<td>Cord, INFOV.A.C.™ China Power</td>
<td>Electri-Cord Mfg. Co.</td>
<td>2.51 m</td>
</tr>
<tr>
<td>4103847</td>
<td>Cord, INFOV.A.C.™ India Power</td>
<td>Electri-Cord Mfg. Co.</td>
<td>2.54 m</td>
</tr>
<tr>
<td>4103887</td>
<td>Cord, Power, Brazil</td>
<td>Electri-Cord Mfg. Co.</td>
<td>2.05 m</td>
</tr>
<tr>
<td>350084</td>
<td>Cord, V.A.C.ULTA ™ AC Power</td>
<td>Various</td>
<td>2.0 m</td>
</tr>
<tr>
<td>350753</td>
<td>Cord, V.A.C.ULTA ™ Power UK</td>
<td>Various</td>
<td>2.0 m</td>
</tr>
<tr>
<td>360074</td>
<td>Cord, V.A.C.VIA ™ Power, EU</td>
<td>Various</td>
<td>2.05 m</td>
</tr>
<tr>
<td>360076</td>
<td>Cord, V.A.C.VIA ™ Power, AU/NZ</td>
<td>Various</td>
<td>2.05 m</td>
</tr>
<tr>
<td>414655</td>
<td>Assy, Power Cord - ACTIV.A.C. ™ Japan</td>
<td>Various</td>
<td>.91 m - 2.44 m (1.0 m preferred)</td>
</tr>
<tr>
<td>415568</td>
<td>KR Cordset</td>
<td>Consolidated Electronics</td>
<td>2.05 m</td>
</tr>
</tbody>
</table>

The use of electrical cables and accessories other than those specified in the supplied instructions for use (IFU) or referenced documents may result in increased electromagnetic emissions from the INFOV.A.C.™ Therapy Unit or decreased electromagnetic immunity of the INFOV.A.C.™ Therapy Unit.
Specifications

Specifications subject to change without notice.

Classification
Equipment not suitable for use in the presence of a flammable anaesthetic mixture with air, oxygen or nitrous oxide.

INFOV.A.C.™ Therapy Unit
Continuous Operation
Type B Equipment
Class I or Internally Powered

Power Supply
Class I Equipment
Ordinary Equipment

INFOV.A.C. Therapy Unit
Dimensions ................................................................. 23 cm X 22 cm X 17.5 cm (9.05 in X 8.6 in X 6.8 in)
Weight .............................................................................. 2.89 kg (6.37 lb)

Electrical Data (Power Supply)
Voltage ........................................................................... 100 - 240 VAC
Frequency .......................................................... 50Hz - 60Hz
Power ............................................................................. 60W

Environmental Conditions
Storage / Transport Temperature Range ................................................................. -10°C to +50°C (14°F to 122°F)
Operational Temperature Range ............................................................................. +10°C to +30°C (50°F to 86°F)
Relative Humidity Range ..................................................................................... 10% to 85% non-condensing
Barometric Pressure Range .................................................................................. 700 hPa to 1060 hPa

The disposable components of the V.A.C.* Therapy System are considered Applied Parts under IEC 60601-1 Third Edition.
Explanation of Symbols Used

⚠️ Warning or Caution statement of possible hazard to system, patient or staff.

ℹ️ Important operational information

⚠️ Caution: Consult Accompanying Documents

⚠️ Tripping Hazard

Power the device on or off. This button will light up when the device is on.

Release the canister. This button will flash when the canister is full and needs to be replaced.

❌ This product is designated for separate collection at an appropriate collection point. Do not dispose of as household waste.

👨‍💻 Refer to User Manual

Authorized Representative in the European Community

Manufacturer

Serial Number
**IPX0**

**Ingress Protection**

**Keep Dry**

**Type B Applied Part**

**Catalog Number**

**No Bathing or Showering**

**Temperature Limits**

**Rx Only**

CAUTION: Federal (US) law restricts this device to sale / rental by or on the order of a physician.

---

**Customer Contact Information**

For questions regarding this product, supplies, maintenance, or additional information about KCI products and services, please contact KCI or a KCI authorized representative, or:

**In the US** call 1-800-275-4524 or visit www.acelity.com.
KCI USA, Inc. 12930 IH 10 West, San Antonio, TX 78249
