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Introduction

VivereX(+)™ system is a medical therapeutic device which applies controlled negative pressure to the wound bed. Healthcare professionals across various disciplines such as physicians, podiatrists, wound specialists, nurses, nurse practitioners, and physician assistants are capable of utilizing the VivereX (+) system. Nevertheless, prior to autonomously utilizing this system, it is imperative for individuals to undergo proper training and attain a requisite level of expertise. The device is equipped with LED indicators for monitoring and alarm the status of the treatment. VivereX(+)™ also has a pressure senso programmed to continuously monitor and maintain a target pressure at the wound site. VivereX™ device is intended to be used for 10 days and the VivereX+™ device for 21 days for one patient, which means no risk for cross extended to the Application of the Application with the Application and the Application applying negative pressure results in transferring of wound exudate to the NPWT dressing. VivereX(+)™ system increases dramatically the dressing life time through its multi-layered design. The VivereX™ dressing anages the wound exudate by two different mechanisms:

The absorbent layer of the dressing absorbs some of the secretion.

- The description of the rest of the wound secretion.

1. Description:

VivereX(+)™ system consists of a single use device. 2 x AA Alkaline batteries viveled(†) System consists of a single use device, 2 A FA mainter settlements fixation strips, an extension tube and VivereX or dressings. The VivereX(+)™ set includes 1 pump unit and 2 NPWT dressings. A multipack of 5 NPWT dressings are available in case additional dressings are required to handle necessary amount of wound exudate. The VivereX™ dressing and VivaFoam are available in following sizes:

NPWT Dressing size/ cm		VivaFoam™ size/ cm		
10 x 20	16 x 16	10 x 7,5 x 3,2	25 x 15 x 1	
10 x 30	16 x 21	20 x 10 x 3,2	15 x 10 x 1	
10 x 40	16 x 31	25 x 15 x 3,2		
21 x 21	26 x 26			

2. Intended use /Indications for use:

VivereX(+) is a novel and multifaceted therapeutic system that facilitates the enhancement of wound healing by utilizing sub-atmospheric pressure to mitigate inflammatory exudate and stimulate the formation of granulation tissue. VivereX(+)^m system is suitable for use in both hospital and home care setting. VivereX(+) system is suitable to be used in adaults, children and neonatals. Following wound types with low and medium exudate are appropriate:

5. Diabetics ulcers

8. Trauma wounds

6. Surgical clean closed incisions
7. Subacute dehisced wounds

9. Burn injury, grade I and II

- Acute wounds
- Chronic wounds
- Skin graft or flaps and donator site 7.
 Pressure ulcers 8.

3. Contraindications:

- VivereX(+)™ system is contraindicated for:
 1. Wounds with malignancy except in palliative care
 2. Wounds with untreated osteomyelitis
- 3. Wounds with untreated non-enteric fistula
- Wounds with exposed bone, blood vessels, nerves, organs or
- anastomotic site
- 6. Tracheostomy and surgical drains

7. Wounds in patients with known allergy to silicone, nylon or acrylic

4. Instruction to use VivereX(+)™ device:

- 1. Insert two AA batteries into the battery compartment on the back side of the VivereX(+)™ device, in the correct orientation (1). Alkaline batteries are recommended. Do not use rechargeable battery, the device will not work. All indicators flash twice indicating the device is ready to use. Secure the battery compartment cover.

 Twist luer connectors together to connect the dressing tube to
- VivereX(+)™ device (2).
- 3. Use extension tube if a longer distance between dressing and the device is required.
 4. Ensure the tube is free from kinks and not blocking the flow between the
- dressing and VivereX(+)™ device.

 5. Press the power button on the front of the VivereX(+)™ device to start the device. The green indicator start to flash, indicating that the device is on.

 6. The device can be heard functioning while
- contraction of the dressing confirms the start of NPWT.

 7. The leak indicator flashes if the dressing is not
- sealed (refer to section 20 for oubleshooting). VivereX(+)™ will turn off if the leak is not adequately solved within 30 seconds.
- Keep VivereX(+)™ in a place close to the patient, where the patient can easily monitor the status of LED indicators e.g. using the belt holder to hang the device on their pocket, outer garments, or belt. 9. The device will automatically stop
- functioning 10 days after start date in VivereX™ and 21 days in VivereX+™.
- 10. To turn off the device, press and hold the



or block exudate.

11. If you require any assistance in setting up, using or maintaining the VivereX(+)™ system, or if any unexpected alerts or fault are encountered, please contact Sunmedic or your local distributer.

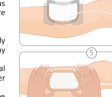
5. Instruction to apply the dressing:

- 1. Using appropriate technique remove hair from area close to the wound in
- order to fix the dressing better to the skin.

 Prepare the wound in accordance to the "Wound Bed Preparation" principals.
- Dry the skin around the wound.
- 3. By the skin around the would.
 4. If the dressing is used on a sensitive skin, wipe the skin around the wound with a supportive solution to protect the skin, prior to the application of the fixation strips. For wounds with moderate volume of secretion.
- the size of the dressing should be approximately 4 times larger than the wound surface. 6. Wounds area with greater than 5 mm in depth should be covered by VivaFoam™ prior to application of the dressing.

 a. Do not Cut VivaFoam™ over the wound as
- fragments may fall into the wound.

 b. Do not force VivaFoam™ into the wound as this might alter the delivery of negative pressure
- Keep in mind that the part of the dressing containing suction port should be placed cranially and outside of the wound over an area of healthy skin (3). 8. Remove the protective layer from the central
- part of the dressing and apply this part over proximal portion of the wound. Remove the rest of the protective layer from the
- dressing and place the dressing on to the wound (4).



- 10. Smooth the dressing with your two hands or pull
- out the dressing edges to prevent creasing (5).

 11. Reposition the dressing if required to avoid crease.

 12. Connect the dressing to the device using luer connectors (2). Use extension tube if it is required.
- 13. Use fixation strips to support and maintain an ose inaction strips to support and maintain an adequate sealed dressing. Apply the fixation strips with a 1cm (0.4 in.) overlap to all sides of the dressing in order to secure a seal

dressing during the wear time (6).

6. Battery change:

The unit is supplied with 2 AA batteries which should provide power to the unit for the period of its lifespan. If the green normal therapy and yellow battery indicators flash, the batteries should be replaced with new ones within 24 hours. If the vellow battery indicator is flashing alone, the battery capacity is too low and the batteries should be replaced within a few hours, preferably immediately

7. Disposal:

Used dressing and fillers should be disposed of as clinical waste in accordance with local hospital protocols. The batteries should be removed from used VivereX(+)TM device and recycled. The device should either be recycled after decontamination or disposed as clinical waste in accordance with the local hospital protocols.

8. Important issues that patients should be advised about:

- 1. Keep the device in a position that allows good visibility of the four LED
 - indicators. 2. Observe routinely tubing to ensure the tubes are free from kinks and not blocking
 - the flow between the dressing and the device.

 3. Keep the device free from dust, drop down.

 - Do not spray or submerge in water.
 Contact your healthcare professional if you observe any evidence of damage or sign of malfunction of VivereX(+)™ system.

9. Taking shower with VivereX™ dressing:

- Press and hold the power button for 3s to put the device in "Off" mode.
 Disconnect VivereX(+)™ and keep the device in a safe place.
- 3. The end of the dressing's tube should be facing down, to prevent water entering the tube.

 4. The dressing should not be submerged in water or sprayed.
- 5. After showering reconnect the device as described in section 4. 6. Press power button to turn the device to "On" mode.
- 7. Ensure that the green indicator flashes, the dressing is contracted and no vellow indicator is active.

10. Cleaning VivereX(+)™ device:

The pump can be cleaned with a damp cloth with soapy water or a gentle disinfectant solution. Avoid spilling liquid on the device as liquids can cause malfunction of VivereX(+)™

device and possibly be a safety hazard.

11. Warnings:

- 1. Treatment with the VivereX(+)™ system is allowed in conjunction with the use of anticoagulants, but because of an increased risk of bleeding, these patients should be monitored continuously, preferably, in a health center.
- 2. In all patients, hemostasis should be done and hemorrhage should be monitored continuously during treatment. If bleeding occurs during treatment with the VivereX(+)™ system, the device should be stopped immediately and promptly notify responsible physician.
- 3. Sharp edges of the bone should be removed or covered in order to avoid
- damage to the veins or internal organs, leading to bleeding.

 4. To minimize the risk of bleeding, remove the dressing carefully.
- Ensure that the dressing is not in direct contact with vessels, nerves, organs or anastomotic sites. These types of tissue should be covered by natural tissue or non-adhesive dressing.
- non-adhesive dressing.

 6. Ensure that the dressing is not placed close to Sympathetic nerve. 6/15

- In case of pediatric patients the size and weight should be taken into consideration.
- 8. Do not use the dressing if the package is damaged.
 9. Disconnect the device if defibrillation is required. Remove the dressing if its
- location interferes with defibrillation. 10. VivereX(+)™ device is not MR compatible and should not be taken into MR
- VivereX(+)[™] device is not compatible with the hyperbaric oxygen environment
- 12. VivereX(+)™ device should not be used in the presence of flammable anesthetic
- 13. Care should be taken to carry device in a way that it does not cause damage,
- or tourniquet or strangulation to the natient Care should be taken that the tubing is not twisted under the patient's body as this may block therapy.
- 15. Keep out of reach of children. Swallowing can lead to chemical burns. perforation of soft tissue, and death. Severe burns can occur within 2 hours of ingestion. Seek medical attention immediately.

12. When to change the dressing:

- A common wear time for a silicone dressing is three to seven days. This period of time reduces in case of wound infection or excessive wound secretion. The dressing should be changed in accordance with following guidelines:
- When the "change dressing" indicator is turned "On" When the wound exudate has reached or is close to the suction port as in (7b) and (7c).
- · When there are repeated sign of leak

13. How to change the dressing:

1. Press and hold the power button for 3s, to turn the device to "Off" mode



- Stretch the fixation strips away from the skin.
 Lift the dressing and remove it with caution horizontally from the surface of the wound.
- 5. Observe the dressing for wound exudate volume, color and odor. Unserve the dressing for wound extudate volume, color and odor.
 Wash the wound with an appropriate wound cleanser if necessary.
 Apply a new dressing as described in section 5.

2. Wait until the dressing is back to its original state.

nnect to the device and press the power button to restart the therapy.

14. To change VivereX(+)™ device:

The VivereX™ has a lifespan of 10 days and the VivereX™ + has a lifespan of 21 days. If the "Leakage", "dressing change" and "Replace Battery" lit continuously, it indicates that the VivereX(+)™ lifespan has elapsed. Press the power button for 3s. Disconnect the device from the unit and connect a new device as described in section 4.

15. Precautions:

- 1. Do not cover the dressing with layers of adhesive film because it reduces the
- evaporation of the wounds secretion and decrease the dressings life time.

 2. Support the fragile skin closed to the wound with skin protecting material.

 3. The VivereX(+)™ pump should not be used with other NPWT accessories other than those mentioned in section 17.
- If the dressing covers circumferentially the extremity monitor the distal circulation carefully and if any sign of ischemia is present, remove the dressing.
- Discontinue using VivereX(+)™ system in case of allergic reaction to the dressing. Infected wounds should be treated by appropriate infection treatment and frequent wound inspections.
- Do not take VivereX(+)™ device apart or modify.
- Any surgical drain should be placed away from the dressing.
 Do not cut the dressing as this cause malfunction. 10. Place the dressing in a way that the suction port of the dressing will be located
- on a healthy skin, away from the wound and cranial to it.

 11. VivereX(+)™ system is a single use device and should not be used on more than
- 12. The dressing and the device should not be exposed to sunshing/or5heat.

13. Do not use the device if the battery cover is missing

- Disconnect the device from the dressing when taking shower. Do not submerge the dressing into water.
- 15. The time needed for minimum and maximum transport/storage temperature to be converted to range of the operational temperature will be approximately
- 16. X-ray might affect some medical devices. If possible, disconnect and move the device out of the X-ray range. Check the device is functioning correctly, If it has

16. Alerts and faults:

VivereX(+)[™] device is equipped with visual indicators to make patient aware of if there is an issue. Refer to section 20, if your device develops a fault. VivereX $(+)^{TM}$ system does not emit audible alarm and the device should be carried in a way that the visual indicators can be monitored easily.

17. Required components for VivereX(+)™ system:

- .. VivereX(+)™ Pump unit
- 2. AA Alkaline battery x 2
- Instructions for use . Sterile dressing
- 5. Extension tube, sterile adhesive fixation strips which are supplied with dressing

18. Caution:

These instructions for use are not intended as a guarantee or warranty. They These instructions for use are not intended as a guarantee of warranty. They are only intended as a guide. For medical questions, please consult a physician This product must be used in accordance with these instructions for use and related labeling

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19. Glossary symbols for VivereX(+)™ device:

①	ON/OFF	~	Device is operating correctly	SN	Serial number
G.	Alert leak through dressing	٦>٠	Dressing need to be changed	③	Follow instruction for use
	Battery alert	\triangle	General warning sign	UDI	Unique Device Identifier
[]i	Consult instructions for use	Ω	Expiry date	REF	Catalogue number
C€	CE mark	®	Do not use if package is damaged	LOT	Batch code
NCN STEPULE	Non-sterile	STERILE EO	Sterilized using ethylene oxide	类	Keep away from sunlight
EC REP	Authorized representative in the European Community	8	Do not reuse	MR	MR unsafe
444	Manufacturer	쎈	Date of manufacture	Lantx	Latex free
75N	Humidity limitation	700 mbar	Atmospheric pressure limitation		Risk of explosion
\$ CO A T T	Temperature limit	P _k only	Prescription use only	Ť	Keep dry
∱	Type BF applied part	A	Not for general waste	I	Fragile handle with care
STEER STEER	Do not resterilize		Do not drop	MD	Medical Device
	Importer		Distibutor		Patient information website

20. Troubleshooting:

Batteries are inserted in

NPWT is not established.

The battery power is low

significant leak exists

NPWT is not established

Possible cause(s)

- The dressing is saturated.

	All indicators are off.	
Possible cause(s)	Troubleshooting/Comments	
The unit is turned off. The batteries are inserted in wrong orientation. The batteries have depleted.	Press the power button to start therapy. Check if the batteries are placed in correct direction. Replace batteries and press power button.	
	All indicators flash twice.	
Possible cause(s)	Troubleshooting/Comments	

The unit is functioning normal. correct orientation Press power button to start therapy The unit is ready for therapy. Possible cause(s) Troubleshooting/Comments The unit is frequently heard running The unit is functioning to maintain the target pressure at the wound hed

· Possible cause(s) Troubleshooting/Comments The air leak should be addressed as

Replace all batteries.

The unit will be turn off after 30s. if

NPWT has not been established due to

air leak. Check dressing's border and smooth out if any creasing is

Troubleshooting/Comments

1- Health care giver to change the

Possible cause(s) Troubleshooting/Comments

observed. Check and secure connectors

?- The tube is blocked. dressing.
2- Ensure the tube is not knicked. 3- The battery power is low

Troubleshooting/Comments

- The dressing is saturated. - Health care giver to change the 2- The tube is blocked. dressing (refer to section 13). 2- Ensure that the tube is not

22. Electromagnetic compatibility:

equipment IEC 60601-1-2.

Possible cause(s)

The unit is functioning normal but the battery power is low.

Possible cause(s)

The unit is functioning normal

but the battery power is too

y **60 (0)**

Possible cause(s)

21. Safety and electromagnetic compatibility:

The VivereX(+)™ Negative Pressure Wound Therapy System complies

with the general requirements for safety of electrical medical equipment IEC 60601 and electromagnetic safety requirements of electrical medical

The unit is expired.

Troubleshooting/Comments

Replace the batteries in less than 24 hours.

he replace battery is flashing

Troubleshooting/Comments

Replace the hatteries within few

Press power button to restart the

hours, preferably immediately.

Troubleshooting/Comments

Health care giver should apply a

if continued therapy is indicated.

new unit and change the dressing

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The therapy is ceased.

VivereX(+)™ Has been tested and shown to comply with the limits for medical to IEC 60601-1-2-2014 4th edition. These limits and test levels are intended to provide reasonable protection to assure the safety of medical devices with regard to electromagnetic disturbances when VivereX(+)™ is used in a typical medical installation and uncontrolled environment like home use. This equipment can be affected by radio frequency energy and if not installed and used in accordance with the instructions, may cause interference to other devices in the vicinity. However, there is no guarantee that interference will not occur in a particular installation. If further information or guidance on electromagnetic immunity and emissions is needed, please contact Sunmedic or visit us online at www.sunmedic.se. The VivereX(+)TM Negative Pressure Wound Therapy system complies with the Essential Performance requirements of IEC 60601- Clause 4.3., to achieve its intended use to maintain a nominal negative pressure of <120 mmHg. VivereX(+)[™] system set (PDI600111-8, PDI600121-8, PDI600411-8, and PDI600421-8) packaged by Sunmedic AB at Hammargatan 11B, 235 32 Vellinge, Sweder

VivereX(+)™ pumps (PDI600100-400) Sunmedic AB Hammargatan 11B, 235 32 Vellinge, Sweden

VivereX[™] FC dressings (PDI600221-8), and VivaFoam[™] (PDI600301-5) Sunmedic AB Hammargatan 11B, 235 32 Vellinge, Sweden

VivereX™ SC dressings (PDI600211-8)
Zhejiang Longterm Medical Technology Co., Ltd.
ShuangShan Road 277, Fuxi Street, Deging County
313200 Huzhou City, Zhejiang Province, China

Lotus NL B.V. Koningin Julianaplein 10, 1e Verd 2595AA The Hague, The Netherlands

Importer and Distributor Sunmedic AB, Hammargatan 11B 235 32 Vellinge, Sweden

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23. Technical Specifications:

Unit weight (excluding batteries)	<100 g		
Dimensions	Height: 84 mm width: 73 mm Thickness: 22 mm		
	ata (Power Supply)		
Power battery	3 V DC/battery		
Input current	Max 300 mA		
Battery type	2 x AA, 2 x 1.5 V, non-rechargeable		
Vacuum mode	Continuous		
Maximum vacuum	120 mmHg		
Operating time, once batteries	10 days for VivereX		
installed	21 days for VivereX+		
Ingress protection	IP22		
Patient protection	Type BF		
Mobility: Portable	Carring clips included/ Carring bag available		
Operation environment	5-40 °C (41-104 °F), 10-75% RH		
	700–1060 mbar atmospheric pressure		
Storage/transport temperature	5-25 °C (41-77 °F), 10-75% RH		
Storage/transport temperature	-20-45 °C (-4-113 °F) in one week		
Altitude range	< 2000 m		
Co	Pump: Non-sterile		
Sterility	Dressing & VivaFoam™: Sterile		
Programable Electronic Medical	SNPC1		
System (PEMS) identifier			
Material Pump (molded parts)	Polycarbonate / ABS		
Material tube	Polyurethane / TPE		
Material dressing	Polyurethane, silicone, absorbent		
Material VivaFoam™	Reticulated open cell polyurethane foam		
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