

# SNAP™ THERAPY SYSTEM

## INSTRUCTIONS FOR USE



Acility®

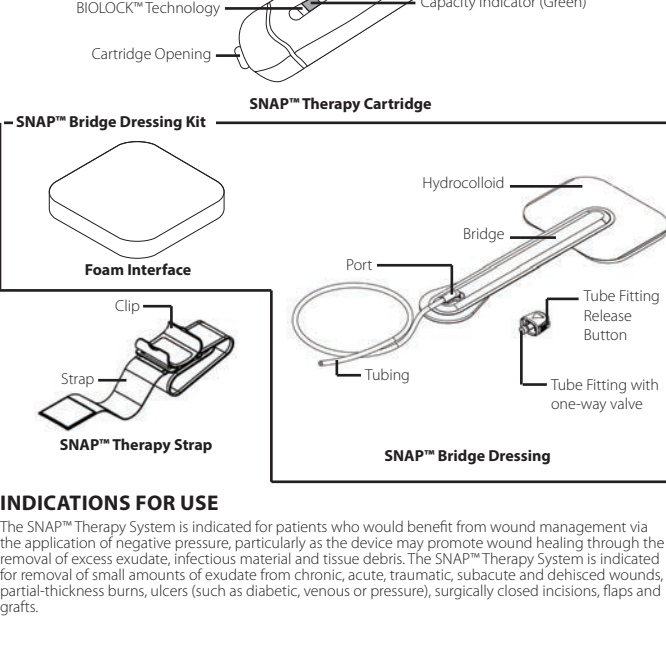
ENGLISH

### SYSTEM DESCRIPTION

The SNAP™ Therapy System is a non-porous, disposable device for the application of negative pressure and a dressing kit for medical dressings. The SNAP™ Bridge Dressing Kit may be suitable for the treatment of acute, traumatic, subacute and chronic wounds, such as wounds on the planter surface of the foot.

Not made with natural rubber latex.

### SNAP™ THERAPY SYSTEM FEATURES

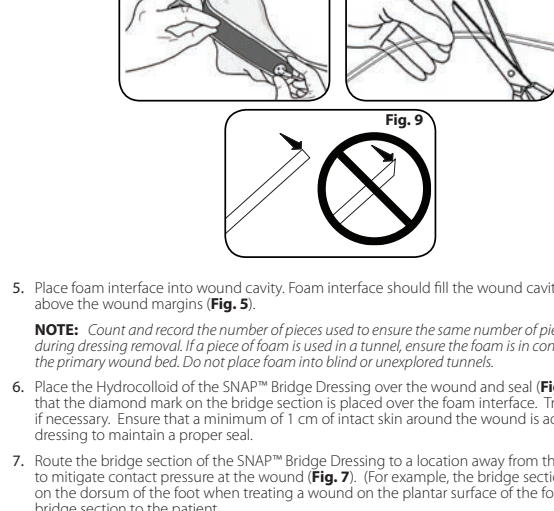


### INDICATIONS FOR USE

The SNAP™ Therapy System is indicated for patients who would benefit from wound management via the application of negative pressure, particularly as the device may promote wound healing through the removal of excess exudate, infectious material and tissue debris.

The SNAP™ Therapy System is indicated for the removal of small amounts of exudate from acute, traumatic, subacute and chronic wounds, partial-thickness burns, burns caused by diabetic, venous or pressure, surgically created incisions, and grafts.

### SYSTEM APPLICATION INSTRUCTIONS (CONT.)



5. Place foam interface into wound cavity. Foam interface should fit the wound cavity and extend over the wound margin [Fig. 5].

**NOTE:** Count and record the number of pieces used to ensure the same number of pieces are removed during dressing removal. If a piece of foam is used in a tunnel, ensure the foam is in contact with foam in the previous tunnel [Fig. 6].

6. Place the Hydrocolloid of the SNAP™ Bridge Dressing over the wound and seal [Fig. 7]. Ensure that the diamond mark on the bridge section is placed over the foam interface. Trim hydrocolloid if excess remains.

7. Route the bridge section of the SNAP™ Bridge Dressing over the wound, location from the toe to the heel of the foot [Fig. 8]. (For example, the bridge section may be placed on the dorsum of the foot when creating a wound on the plantar surface of the foot). Secure the bridge to the patient.

8. Cut the dressing tubing to the desired length [Fig. 8].

**NOTE:** The tube should be straight, not an angle. [Fig. 9]. This will give a proper seal when connected to the tube fitting.

9. Fully insert the tube fitting into the tubing [Fig. 10].



10. Connect the SNAP™ Therapy Cartridge, press down on activation / reset key and pull it out [Fig. 11]. Re-insert and repeat as needed until an airtight seal is obtained and the red pressure discharge indicator is not visible in the pressure discharge window.

11. To activate the SNAP™ Therapy Cartridge, press down on activation / reset key and pull it out [Fig. 12]. Re-insert and repeat as needed until an airtight seal is obtained and the red pressure discharge indicator is not visible in the pressure discharge window.

12. Secure the SNAP™ Therapy Cartridge to the patient's extremity or belt using the SNAP™ Therapy Strap [Fig. 13].

**NOTE:** When the strap is placed around an extremity, take care to ensure that the strap is not placed too tightly around the extremity. This may result in a decrease in blood flow to the extremity. Digital perfusion may be assessed by noting skin color, altered sensation or pulses.

13. Check negative pressure operating. The SNAP™ Therapy System is working properly if:

- Green capacity indicator is both visible and stationary in the chamber window.
- Dressing has a suction hold down appearance.
- The dressing feels hard to the touch.

Regular visual inspection of the SNAP™ Therapy System is recommended so that any loss in negative pressure delivery can be recognized in a timely manner.

**NOTE:** At a minimum the SNAP™ Therapy Cartridge should be inspected once every eight hours. It is recommended that the dressing be changed at a minimum of two times per week, with frequency dictated by the physician or practitioner.

14. Cut the dressing tubing to the desired length [Fig. 8].

**NOTE:** The tube should be straight, not an angle. [Fig. 9]. This will give a proper seal when connected to the tube fitting.

15. Fully insert the tube fitting into the tubing [Fig. 10].



16. Connect the SNAP™ Therapy Cartridge, press down on activation / reset key and pull it out [Fig. 11]. Re-insert and repeat as needed until an airtight seal is obtained and the red pressure discharge indicator becomes visible before an airtight seal is established. Reset the SNAP™ Therapy Cartridge using the following steps:

17. Turn the cartridge counter-clockwise to remove the tube fitting from the tube fitting.

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# SNAP™-BEHANDLINGSSYSTEM

BRUKSANVISNING  
SNAP™-behandlingspatron  
SNAP™-broforbindingssett  
SNAP™-behandlingsstropp



NORGK

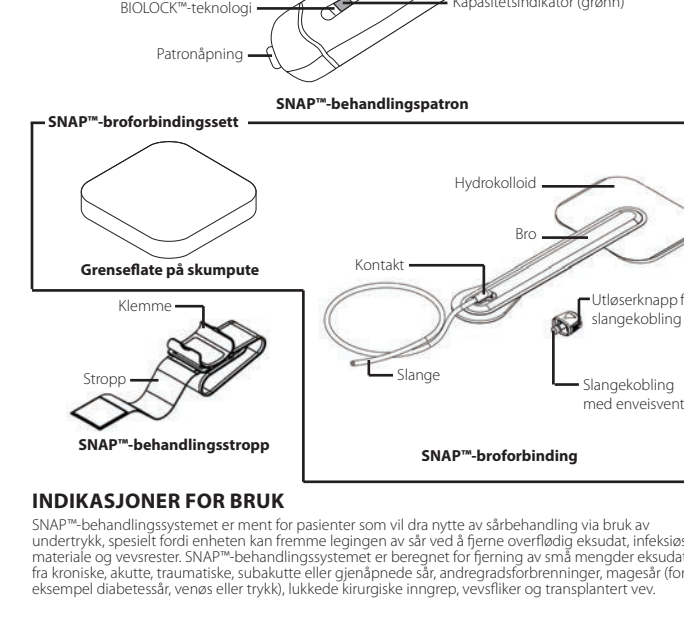


## SYSTEMBESKRIVELSE

SNAP™-behandlingsystemet omfatter et ikke-motoriseret engangsinnlegg for anvendelse av undertrykk og et forbindingssett for medisinsk bruk. SNAP™-broforbindingssettet kan være egnet for slapsplasering som kan ha myte av å posisjonere patienten bort fra sletten, for eksempel så på fotens plantaroverflate.

Ikke laget med naturgummimateriale.

## FUNKSJONER I SNAP™-BEHANDLINGSSYSTEMET



## INDIKASJONER FOR BRUK

SNAP™-behandlingspatronen kan brukes for pasienter som vil få nytte av tilberedning via bruk av undertrykk, spesielt før enheten kan fremme leggningen av slett ved å fjerne overflaten eksudat, infeksjon, materiale og vevrestre. SNAP™-behandlingssystemet er beregnet for fremming av små mängder eksudat fra kontrollert og kontinuerlig utslipp av eksudat, og kan redusere risikoen for infeksjoner, blodutløp og tørkeproblemer.

Ikke tilpasset for bruk i føtene til pasienter med diabetes, venær eller trykki, lukkede kinskrueinnlegg, vevrestre og transmetten ved.

## INSTRUKSJONER FOR BRUK (FORTS.)



5. Passer skumoverflaten i såhulen. Skumoverflaten skal fylle såhulen og strekke seg over sårkontakten. **Fig. 5**

MERK: Tell og registrer entlastede deler som henviser til at ikke det samme antastes deler ferner når forbindingen fjernes. Hvis en bit av skumputen brukes i en gang, må du kontrollere at skumputen er i kontakt med skumputen i den primære såhulen.

6. Passer hydrokloridet til SNAP™-broforbindingen over slett og føres. **Fig. 6** Sørg for at diamantmerket på brodenken er plassert over skumoverflaten. Klipp om nødvendig til hydrokloridet ordentlig førstegang.

7. Plissert enden av SNAP™-behandlingsstroppen et sted innan slett for å redusere kontaktkraft på slett ved behandling av et slett på fotens plantaroverflate. Fest brodenken til patienten. **Fig. 7** (Bemerk: kan for eksempel plasseres på domus av foten ved behandling av et slett på fotens plantaroverflate). Fest brodenken til patienten.

8. Klipp forbindingslangen til ønsket lengde. **Fig. 8**

MERK: Kurvet skal være rett, ikke vinklet. **Fig. 9** Det gir riktig forslagning når slangen holdes til.

Regelmessig vurdering av teknikkene i denne instruksjonen.

Regelmessig vurdering av teknikk

