

## 510(k) Summary

JUN 22 2011

Submitted by: Coreleader Biotech Co., Ltd.  
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Contact Teeming Tsao  
 Person:  
 Date Prepared: September 20, 2010

Proprietary Coreleader AlgiPlaster  
 Name:  
 Common Topical Wound Dressing  
 Name:  
 Classification: Unclassified

Classification Hydrophilic Wound and Burn Dressing  
 Name:  
 Predicate CALGON VESTAL DIV., K910059, KALTOSTAT WOUND  
 Device: DRESSING  
 COLOPLAST CORP., K983519, COMFEEL SEASORB  
 DRESSING

	COMFEEL SEASORB DRESSING	CORELEADER ALGIPLASTER
Device description	The Seasorb Dressing is a highly absorbent wound dressing consisting of an calcium sodium alginate/sodium carboxymethylcellulose Xerogel cast into a high density polyethylene net.	The Coreleader AlgiPlaster is a soft, sterile, non-woven dressing of calcium alginate fibers combined with a thin polyurethane membrane. It provides a barrier to external contamination and produces a moist environment at the surface of the wound

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		by reducing water vapor loss from the exposed tissue. The moist environment provided by the alginate fibers may provide an environment favorable to the wound healing process.
Use(single, reusable)	Single	Single
Bio-compatibility	Non-hypersensitivity · non-cytotoxicity and non-irritant	Non-hypersensitivity · non-cytotoxicity and non-irritant
Technological features	Calcium alginate fibers are based heavily upon its unique gelation properties with respect to mono- and divalent cations.	Calcium alginate fibers are based heavily upon its unique gelation properties with respect to mono- and divalent cations.
Intended use	For management of (under the guidance of a health care professional) moderate to heavily exudating wounds, including leg ulcers and pressure sores, etc.	The following conditions are considered appropriate for OTC use by the lay person: Coreleader AlgiPlaster is indicated for the management minor abrasions, minor cuts and minor lacerations.  The following conditions are considered appropriate for Prescription Use Under the supervision of a licensed healthcare

		<p>provider: Coreleader AlgiPlaster is indicated as a primary dressing for the management of exuding wounds, including acute wounds such as abrasions, lacerations and post-surgical wounds.</p>
<p>Precautions</p>	<p>Wounds which are solely or mainly caused by arterial insufficiency or complicated diabetic wounds (primarily lower leg and foot) should be inspected by a physician or nurse regularly. A physician should be consulted before using this product on wound with a high risk of infection, or on lesions caused by syphilis, tuberculosis, leprosy or cancer. Comfeel Seasorb dressing must be removed prior to the following treatments: radiation, X-rays, ultrasonic treatment, diathermy and micro waves. Wounds with signs of clinical infection, fever</p>	<p>Rx precautions: The Wound Dressing may be used on high risk of infected wounds only under the care of clinical physicians. AlgiPlaster must not be used as a surgical implant. The dressing can't be left in the wound permanently and should be inspected by a physician or nurse regularly. Wounds with signs of clinical infection, fever and local symptoms such as pain, erythema or pus should have a bacterial swab examination. Use of this product may be continued at the discretion of a physician. Current systemic antibiotic treatment may be give if indicated. The product is for single use only and should not be re-sterilized. Reuse or</p>

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	<p>and local symptoms such as pain, erythema or pus should have a bacterial swab examination. Use of this product may be continued at the discretion of a physician. Current systemic antibiotic treatment may be give if indicated.</p> <p>Not recommended for use on dry wounds or third degree burns.</p> <p>Do not use on patients with known hypersensitivity to any of the ingredients.</p>	<p>re-sterilization may also create a risk of contamination of the device and/or cause patient infection or cross-infection.</p> <p>Not recommended for use on dry wounds or third degree burns.</p> <p>Do not use if the package is opened or the seal broken.</p> <p>OTC precautions: Do not use on infected wounds</p> <p>The dressing can't be left in the wound permanently.</p> <p>Reuse or re-sterilization may also create a risk of contamination of the device and/or cause patient infection or cross-infection.</p> <p>The product is for single use only and should not be re-sterilized.</p> <p>Not recommended for use on dry wounds or third degree burns.</p> <p>Do not use if the package is opened or the seal broken.</p> <p>Do not use beyond the expiration date.</p>
Sterilization	Sterile	Sterile
Packaging	Polyester pouches laminated with peelable	Sterilization pouch: Top material: Medical

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	polyethylene prior to sterilization	grade paper pouches. Bottom material: Transparent see-through film,(made of PET).
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**Device Description:** Coreleader AlgiPlaster is composed of Alginate fiber and combined with the thin polyurethane membrane coated with a layer of an acrylic adhesive. The dressing combination, which is permeable to both water vapor and oxygen, is impermeable to micro-organisms and once in position, it provides an effective barrier to external contamination, and producing a moist environment at the surface of the wound by reducing water vapor loss from the exposed tissue. The moist environment provided by the alginate fibers may provide an environment favorable to the wound healing process.

Coreleader AlgiPlaster is a sterile topical wound dressing, packed in individual pouch and sterilized by r-ray radiation to a  $10^{-6}$  SAL.

**Intended Use:** The following conditions are considered appropriate for OTC use by the lay person:

Coreleader AlgiPlaster is indicated for the management minor abrasions, minor cuts and minor lacerations.

The following conditions are considered appropriate for Prescription Use under the supervision of a licensed healthcare provider:

Coreleader AlgiPlaster is indicated as a primary dressing for the management of exuding wounds, including acute wounds such as abrasions, lacerations and post-surgical wounds.

**Technological Characteristics :** Coreleader AlgiPlaster is composed of calcium alginate fiber and PU membrane. The calcium alginate fiber is manufactured by a Wet-Spinning production process. Alginate fiber combined with the thin polyurethane membrane coated with a layer of an acrylic adhesive. The dressing combination, which is permeable to both water vapour and oxygen, is impermeable to micro-organisms and once in position, it provides an effective barrier to external contamination, and producing a moist environment at the surface of the wound by reducing water vapour loss from the exposed tissue. The dressing is presented as a dense, flat non-woven pad for

application to surface wounds. In the presence of exudate or other body fluids containing sodium ions, the fibers absorb liquid and swell. Calcium ions present in the fibers are partially replaced by sodium ions, causing the dressing to take on a gel-like consistency which may provide an environment favorable to the wound healing process.



Food and Drug Administration  
10903 New Hampshire Avenue  
Document Control Room -WO66-G609  
Silver Spring, MD 20993-0002

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% Mr. Ian Li  
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Sintai 5<sup>th</sup> Rd., Sijhih City  
Taipei, Taiwan 22102

JUN 22 2011

Re: K102942  
Trade/Device Name: Coreleader AlgiPlaster  
Regulatory Class: Unclassified  
Product Code: FRO  
Dated: May 20, 2011  
Received: June 07, 2011

Dear Mr. Li:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you; however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21

CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

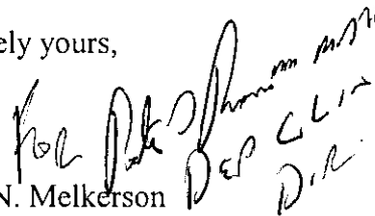
If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please go to <http://www.fda.gov/AboutFDA/CentersOffices/CDRH/CDRHOffices/ucm115809.htm> for the Center for Devices and Radiological Health's (CDRH's) Office of Compliance. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

<http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>.

Sincerely yours,

Handwritten signature of Mark N. Melkerson in black ink. The signature is stylized and includes the initials 'M.N.M.' and 'D.R.'.

Mark N. Melkerson  
Director  
Division of Surgical, Orthopedic  
And Restorative Devices  
Office of Device Evaluation  
Center for Devices and  
Radiological Health

Enclosure

## Indications for Use Statement

510(k) Number (if known): K102942

Device Name: Coreleader AlgiPlaster

### Indications for Use:

The following conditions are considered appropriate for OTC use by the lay person: Coreleader AlgiPlaster is indicated for the management minor abrasions, minor cuts and minor lacerations.

The following conditions are considered appropriate for Prescription Use under the supervision of a licensed healthcare provider:

Coreleader AlgiPlaster is indicated as a primary dressing for the management of exuding wounds, including acute wounds such as abrasions, lacerations and post-surgical wounds.

Prescription Use   V    
(Part 21 CFR 801 Subpart D)

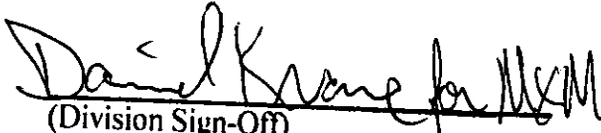
AND/OR

Over-The-Counter Use   V    
(21 CFR 801 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

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Concurrence of CDRH, Office of Device Evaluation (ODE)

  
(Division Sign-Off)  
Division of Surgical, Orthopedic,  
and Restorative Devices

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