

Federal Employee Program® 1310 G Street, N.W. Washington, D.C. 20005 202.942.1000 Fax 202.942.1125

# 5.90.031

Section: Prescription Drugs Effective Date: October 1, 2023

Subsection: Topical Products Original Policy Date: May 12, 2017

Subject: Santyl Page: 1 of 4

Last Review Date: September 8, 2023

# Santyl

#### **Description**

Santyl (collagenase)

#### **Background**

Santyl (collagenase) ointment is a sterile enzymatic debriding ointment which contains 250 collagenase units per gram of white petrolatum USP. The enzyme collagenase is derived from the fermentation by *Clostridium histolyticum*. It possesses the unique ability to digest collagen in necrotic tissue (1).

#### **Regulatory Status**

FDA-approved indication: Santyl ointment is indicated for debriding chronic dermal ulcers and severely burned areas (1).

#### Related policies

Regranex

### **Policy**

This policy statement applies to clinical review performed for pre-service (Prior Approval, Precertification, Advanced Benefit Determination, etc.) and/or post-service claims.

Santyl ointment may be considered **medically necessary** if the conditions indicated below are met.

Santyl ointment may be considered **investigational** for all other indications.

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### **Prior-Approval Requirements**

#### **Diagnoses**

Patient must have **ONE** of the following:

- 1. Chronic dermal ulcer
- 2. Severely burned areas

#### **AND ALL** of the following:

- a. Documented presence of necrotic tissue, sinus tracts, exudation or infection of soft and hard tissues
- b. Prescriber agrees to terminate treatment when debridement of necrotic tissue is complete and granulation tissue is well established

## Prior – Approval Renewal Requirements

#### **Diagnoses**

Patient must have **ONE** of the following:

- 1. Chronic dermal ulcer
- 2. Severely burned areas

#### **AND ALL** of the following:

- a. Improvement in wound
- b. Prescriber agrees to terminate treatment when debridement of necrotic tissue is complete and granulation tissue is well established

### **Policy Guidelines**

#### Pre - PA Allowance

None

# **Prior - Approval Limits**

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**Quantity** 360 grams per 90 days

**Duration** 3 months

# Prior - Approval Renewal Limits

Same as above

#### Rationale

#### **Summary**

Santyl ointment is a sterile enzymatic debriding ointment which contains 250 collagenase units per gram of white petrolatum USP. The enzyme collagenase is derived from the fermentation by *Clostridium histolyticum*. It possesses the unique ability to digest collagen in necrotic tissue. Santyl ointment is indicated for debriding chronic dermal ulcers and severely burned areas. Use of Santyl ointment should be terminated when debridement is complete and granulation tissue is well established (1).

Prior authorization is required to ensure the safe, clinically appropriate, and cost-effective use of Santyl ointment while maintaining optimal therapeutic outcomes.

#### References

1. Collagenase Santyl [package insert]. Fort Worth, Tx. Smith & Nephew, Inc.; 2016.

Policy History	
Date	Action
May 2017	Addition to PA
June 2017	Annual review
June 2017	Update of the tried and failed agents
September 2018	Annual review
March 2019	Removed requirement of inadequate treatment response, intolerance, or contraindication to iodosorb or OTC wound debridement gel or dressing
June 2019	Annual review
September 2020	Annual review
September 2021	Annual review
September 2022	Annual review
September 2023	Annual review

#### Keywords

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This policy was approved by the FEP® Pharmacy and Medical Policy Committee on September 8, 2023 and is effective on October 1, 2023.