

# Healing efficacy, safety and ease of use of a thin Algostérial

<b>Submission date</b> 11/01/2017	<b>Recruitment status</b> No longer recruiting	<input type="checkbox"/> Prospectively registered
<b>Registration date</b> 19/01/2017	<b>Overall study status</b> Completed	<input type="checkbox"/> Protocol
<b>Last Edited</b> 08/08/2018	<b>Condition category</b> Skin and Connective Tissue Diseases	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
		<input type="checkbox"/> Individual participant data
		<input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Patients with open wounds often need to be treated with a dressing to help the wound to heal. Algostérial is a calcium alginate wound dressing, made from seaweed. It works by providing a moistened environment for the wound which helps the healing process. This study is looking at a new form of Algosteril, narrower than Algosteril, commonly used by surgeons. The aim of this study is to find out whether this new shape of Algosteril, which can better adapt to the shape of wounds, can help improve wound healing.

### Who can participate?

Adults with a wound that needs dressing.

### What does the study involve?

Each patient is treated using Algostérial dressings until their wound is healed. The dressing is replaced when required (every two days maximum) until it is no longer required. The length of wound healing is around one month, however this may vary depending on the patient. The wound is examined every time the dressing is changed in order to look for any signs of infection and to record the length of time taken for the wound to heal.

### What are the possible benefits and risks of participating?

The potential benefits to participating in this study include quick wound healing, using a dressing that is easy to use and remove. There are no notable risks involved with participating in this study.

### Where is the study run from?

1. Pitié-Salpêtrière Hospital (France)
2. CHU Estaing (France)
3. Centre François-Xavier Michelet (France)
4. Jean Minjoz Hospital (France)

### When is the study starting and how long is it expected to run for?

February 2016 to December 2017

Who is funding the study?  
Les Laboratoires Brothier (France)

Who is the main contact?  
Dr Mueser Maryse

## Contact information

**Type(s)**  
Scientific

**Contact name**  
Dr Maryse Mueser

**Contact details**  
Les Laboratoires Brothier  
41 rue de Neuilly  
Nanterre  
France  
92735

## Additional identifiers

**Protocol serial number**  
N°ID RCB : 2016-A00101-50

## Study information

**Scientific Title**  
Healing efficacy, safety and ease of use of a thin Algostéril in the wounds of maxillofacial surgery

**Study objectives**  
The aim of the study is to demonstrate the efficiency of thin Algostéril in wounds of the head and the neck.

**Ethics approval required**  
Old ethics approval format

**Ethics approval(s)**  
Persons Protection Committee, CPP Ile-De-France VI, 16/03/2016, ref: CPP/7-16

**Study design**  
Multi-centre prospective non-randomised study

**Primary study design**  
Interventional

**Study type(s)**  
Treatment

## **Health condition(s) or problem(s) studied**

Wound care

## **Interventions**

All patients have their wounds treated using an Algostéril dressing until wound healing (about one month). Wounds will be redressed every two days maximum. Participants are clinically examined in order to determine the number of days until wound healing occurs. There will be no follow up post-healing.

## **Intervention Type**

Other

## **Primary outcome(s)**

Number of days of treatment to obtain the wound healing is assessed through clinical examination until time of wound healing (approximately 1 month).

## **Key secondary outcome(s)**

1. Draining efficiency evaluated by exudate quantity and local signs of infection at the time of redressing (every 2 days maximum)
2. Ease of use is assessed using a scale of 1 to 4 at the time of redressing (every 2 days maximum)
3. Safety is assessed by recording the type and frequency of adverse events continuously throughout the study

## **Completion date**

31/05/2019

# **Eligibility**

## **Key inclusion criteria**

1. Patients with a wound that needs to be treated with thin Algostéril
2. Those who can be followed until the wound healing
3. Signed informed consent form
4. Aged 18 years and over

## **Participant type(s)**

Patient

## **Healthy volunteers allowed**

No

## **Age group**

Adult

## **Lower age limit**

18 years

## **Sex**

All

## **Key exclusion criteria**

1. Pregnant women
2. Participation in another clinical trial within 30 days prior to inclusion

**Date of first enrolment**

16/01/2017

**Date of final enrolment**

30/04/2019

## Locations

**Countries of recruitment**

France

**Study participating centre****Pitié-Salpêtrière Hospital**

47-83 Boulevard de l'Hôpital

Paris

France

75013

**Study participating centre****CHU Estaing**

1 Rue Lucie Aubrac

Clermont-Ferrand

France

63100

**Study participating centre****Centre François-Xavier Michelet**

Place Amélie Raba-Léon

Bordeaux

France

33076

**Study participating centre****Jean Minjoz Hospital**

3 Boulevard A. Fleming

Besançon

France

25030

# Sponsor information

## Organisation

Les Laboratoires Brothier

## ROR

<https://ror.org/007jkh405>

# Funder(s)

## Funder type

Industry

## Funder Name

Les Laboratoires Brothier

# Results and Publications

## Individual participant data (IPD) sharing plan

The current data sharing plans for the current study are unknown and will be made available at a later date.

## IPD sharing plan summary

Data sharing statement to be made available at a later date