

# A study on cord blood stem cells and platelet-rich plasma for treating pregnancy stretch marks

<b>Submission date</b> 01/11/2021	<b>Recruitment status</b> No longer recruiting	<input checked="" type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
<b>Registration date</b> 09/11/2021	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 05/11/2021	<b>Condition category</b> Skin and Connective Tissue Diseases	<input type="checkbox"/> Individual participant data <input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

Background and study aims:

Pregnancy stretch marks (striae gravidarum) are very common among pregnant women and may cause emotional and psychological stress. Current treatments, however, are not satisfactory. As cord blood-derived platelet-rich plasma (PRP) gel and stem cells have good regenerative capabilities, many studies have found the combination works well in cosmetic applications. The aim of this study is to test the effectiveness of PRP stem cell therapy at treating pregnancy stretch marks.

Who can participate?

Pregnant women aged 23 to 30 who have pregnancy stretch marks

What does the study involve?

Participants have their cord blood collected when giving birth. The collected blood provides the PRP and stem cells used later. The left abdomen of each participant receives no treatment. The right abdomen of each participant is treated with PRP stem cell injections on day 1, day 10, and day 20 after giving birth. The research staff will take photos of the abdomen to see how well the treatment works as time progresses.

What are the possible benefits and risks of participating?

The possible benefits include a safe and effective treatment of the stretch marks and tolerance for cell therapy. The risks involve the body reacting badly to the transplantation (it is not very probable since the materials used are from one's own body).

Where is the study run from?

Shenyang Maternal and Child Care Service Center, Shenyang, Liaoning (China)

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

October 2021 to June 2023

Who is funding the study?  
Shenyang Engineering Technology R&D Center of Cell Therapy Co., Ltd. (China)

Who is the main contact?

1. Dr Ning Pei  
775104436@qq.com
2. Prof. Yanqiu Yu  
yqyu@cmu.edu.cn
3. Dr Jing Li  
820367546@qq.com

## Contact information

### Type(s)

Scientific

### Contact name

Dr Ning Pei

### Contact details

No.41 Shenzhou Rd  
Heping District  
Shenyang  
China  
110004  
+86 (0)24 13066597504  
775104436@qq.com

### Type(s)

Scientific

### Contact name

Prof Yanqiu Yu

### Contact details

Shenyang Engineering Technology R&D Center of Cell Therapy Co., Ltd  
A10,400-8 wisdom Second Street  
Hunnan District  
Shenyang  
China  
110004  
+86 (0)24 13386887215  
yqyu@cmu.edu.cn

### Type(s)

Public

### Contact name

Ms Yutong Ge

**Contact details**

Shenyang Engineering Technology R&D Center of Cell Therapy Co., Ltd  
A10,400-8 wisdom Second Street  
Hunnan District  
Shenyang  
China  
110004  
+86 (0)24 15142525031  
yutong@sycellcenter.com

**Type(s)**

Scientific

**Contact name**

Dr Jing Li

**Contact details**

No.41 Shenzhou Rd, Heping District  
Shenyang  
China  
110004  
+86 (0)24 13998160669  
820367546@qq.com

**Additional identifiers****EudraCT/CTIS number**

Nil known

**IRAS number****ClinicalTrials.gov number**

Nil known

**Secondary identifying numbers**

Nil known

**Study information****Scientific Title**

The effect of autologous umbilical cord blood stem cells and platelet-rich plasma on striae gravidarum

**Acronym**

PRPPSM

**Study objectives**

The combination of cord blood-derived autologous platelet-rich plasma (PRP) gel and stem cells is effective in treating striae gravidarum.

**Ethics approval required**

Old ethics approval format

**Ethics approval(s)**

Approved 29/10/2021, Institutional Review Board of the Shenyang Maternal and Child Care Service Center (No.41 Shenzhou Rd, Heping District, Shenyang, China; +86 (024)86906168; 775104436@qq.com), ref: 2021-001-01

**Study design**

Single-center non-randomized study

**Primary study design**

Interventional

**Secondary study design**

Non randomised study

**Study setting(s)**

Hospital

**Study type(s)**

Treatment

**Participant information sheet**

No participant information sheet available

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

Striae gravidarum

**Interventions**

Control group: left abdomen of each participant, no treatment.

Intervention group: right abdomen of each participant, PRP stem cell transplantation.

Transplantation:  $1 \times 10^7$  autologous cord blood stem cells and 15 ml autologous PRP gel, administered via subcutaneous injection at the site of the striae gravidarum. Three transplantations are planned at Day 1, 10, and 20 post-delivery.

Additional procedures:

1. Before the injection:

Apply lidocaine cream at the injection site beforehand to reduce pain caused by the puncture

2. During and immediately after the injection:

1. Apply manual pressure at the injection area for an extended period to reduce bleeding

2. Apply cold compress at the injection area to prevent bruise

3. In case of a bruise, at 48 h post-transplantation:

Apply hot compress at the bruised area to facilitate absorption

**Intervention Type**

Biological/Vaccine

**Phase**

Not Applicable

**Primary outcome measure**

Visual changes measured using abdomen photos at baseline, the 7th day after each transplantation (i.e., day 8, 17, and 27 post-delivery), month 2, and month 3 post-delivery

**Secondary outcome measures**

Measured at baseline, day 8, 17, 27, month 2, and month 3 post-delivery:

1. Pain caused by striae gravidarum measured using self-reporting questionnaires
2. Pruritus caused by striae gravidarum measured using self-reporting questionnaires

**Overall study start date**

01/10/2021

**Completion date**

30/06/2023

## Eligibility

**Key inclusion criteria**

1. Pregnant women with striae gravidarum
2. Full-term or cesarean delivery
3. Aged 23 to 30 years

**Participant type(s)**

Patient

**Age group**

Adult

**Sex**

Female

**Target number of participants**

10

**Key exclusion criteria**

1. Participant in other ongoing clinical trials
2. Tested positive for hepatitis B, hepatitis C, HIV, syphilis, EBV, or CMV
3. History of severe allergic reactions or adverse reactions to medication

**Date of first enrolment**

15/11/2021

**Date of final enrolment**

14/11/2022

## Locations

**Countries of recruitment**

China

**Study participating centre****Shenyang Maternal and Child Care Service Center**

No.41 Shenzhou Rd

Heping District

Shenyang

China

110004

## **Sponsor information**

**Organisation**

Shenyang Maternal and Child Care Service Center

**Sponsor details**

No.41 Shenzhou Rd

Heping District

Shenyang

China

110004

+86 (0)2486906168

syfykj2011@163.com

**Sponsor type**

Hospital/treatment centre

**Website**

<http://sysfybjy.net/index.php?g=&m=Index&a=index>

**Organisation**

Shenyang Engineering Technology R&D Center of Cell Therapy Co., Ltd

**Sponsor details**

A10, 400-8 wisdom Second Street

Hunnan District

Shenyang

China

110004

+86 (0)15142525031

shenyangcell@163.com

**Sponsor type**

Industry

**Website**

<http://www.sycellcenter.com/index.php/cn/Index/index.html>

**Funder(s)****Funder type**

Industry

**Funder Name**

Shenyang Engineering Technology R&D Center of Cell Therapy Co., Ltd

**Results and Publications****Publication and dissemination plan**

Planned publication in a high-impact peer-reviewed journal

**Intention to publish date**

31/12/2022

**Individual participant data (IPD) sharing plan**

The 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