# Effect of oral probiotics on skin wrinkles and hydration in adult women

Submission date 28/09/2020	<b>Recruitment status</b> No longer recruiting	[X] Prospectively registered
		[_] Protocol
Registration date	Overall study status	[] Statistical analysis plan
05/10/2020	Completed	[X] Results
Last Edited 09/01/2025	<b>Condition category</b> Skin and Connective Tissue Diseases	Individual participant data

### Plain English summary of protocol

Background and study aims

Bifidobacteria are bacteria that live in the human gut. They help to improve digestion of food and suppress the growth of harmful bacteria. Gut bacteria may also have other positive effects on human heath. Bifidobacteria and components derived from Bifidobacteria have been studied for their properties in enhancing skin hydration and structure. The aim of this study is to examine the effects of taking Bifidobacterium by mouth on the skin, including wrinkles, dryness and elasticity.

Who can participate? Korean women aged 30-60 years

### What does the study involve?

Participants will be randomly allocated to one of two groups. Both groups will take a capsule by mouth once a day for 12 weeks. For one group, the capsule will contain Bifidobacterium. For the other, the capsule will be a dummy capsule containing no active ingredient (placebo). The capsules will look the same and neither the participants nor the researchers will know which capsule a participant is taking.

What are the possible benefits and risks of participating?

Participants may not benefit from the study as it is not known whether the test product will improve skin wrinkles or hydration. However, the study will extend the understanding of the effects of oral Bifidobacterium on skin wrinkles and hydration.

As for potential risks and disadvantages, blood sampling may cause a hematoma (bruise) or fainting, and the skin measurements and restrictions related to the measurements may cause inconvenience. The test product is not known to cause side effects. There is a small risk of possible allergy or intolerance, as for any dietary product.

Where is the study run from? Danisco Sweeteners Oy (Finland)

When is the study starting and how long is it expected to run for? September 2019 to March 2021 Who is funding the study? Danisco Sweeteners Oy (Finland)

Who is the main contact? Dr Ja Hyun Ryu, dermapro@dermapro.co.kr

### **Contact information**

**Type(s)** Public

**Contact name** Dr Ja Hyun Ryu

### **Contact details**

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### Type(s)

Scientific

**Contact name** Ms Laura Huuskonen

### **Contact details**

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# Additional identifiers

**EudraCT/CTIS number** Nil known

**IRAS number** 

**ClinicalTrials.gov number** Nil known

Secondary identifying numbers

# Study information

### Scientific Title

Effects of Bifidobacterium on skin wrinkle, hydration, ransepidermal water loss (TEWL), elasticity and gloss: a randomized, double-blind, placebo controlled, parallel clinical trial

### Acronym

RM\_StructProb-A

### **Study objectives**

Consumption of a Bifidobacterium-based probiotic reduces skin wrinkles and enhances skin hydration in adult women.

### Ethics approval required

Old ethics approval format

### Ethics approval(s)

Approved 06/08/2020, DERMAPRO Ltd. Institutional Review Board (4F, 30, BangbaeJungang-ro, Seocho-gu, Seoul, Korea; +82-2-597-5435; dermapro@dermapro.co.kr), ref: 1-220777-A-N-01-DICN20181

### **Study design** Randomized double-blind parallel placebo-controlled study

**Primary study design** Interventional

**Secondary study design** Randomised controlled trial

**Study setting(s)** Other

**Study type(s)** Treatment

### Participant information sheet

Not available in web format, please use contact details to request a participant information sheet.

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

Skin wrinkles and dry skin

### Interventions

Participants are randomly allocated to one of two treatment groups in equal proportions applying block randomization.

They receive 1 daily capsule of test product (Bifidobacterium in microcrystalline cellulose) or placebo (microcrystalline cellulose) for 12 weeks with follow-up visits at 4, 8 and 12 weeks.

During the study participants will need to follow some lifestyle restrictions to allow reliable measurements of the selected skin properties. These restrictions include consumption of probiotics, use of certain topical products and exposure to excessive sunlight. The study includes 5 visits to the study clinic. All visits will be carried out at DERMAPRO Ltd., Seoul, Korea. Before each visit, make-up needs to be avoided to assure reliable measurement results.

### Intervention Type

Supplement

### Primary outcome measure

1. Skin wrinkles around crow's feet measured with PRIMOS® imaging technology at baseline and 4, 8 and 12 weeks.

2. Skin hydration measured at cheek, forearm and back of hand with corneometer and moisturemeter at baseline and 4, 8 and 12 weeks.

### Secondary outcome measures

1. Transepidermal water loss from cheek, forearm and back of hand measured using evaporimeter at baseline and 4, 8 and 12 weeks

2. Cheek skin elasticity measured using cutometer at baseline and 4, 8 and 12 weeks

3. Cheek skin gloss measured using SkinGlossMeter at baseline and 4, 8 and 12 weeks

4. Body composition measured with Inbody 330 analyzer at baseline and 2 weeks

5. Facial image measured using VISIA-CR® skin analysis system at baseline and 4, 8 and 12 weeks 6. Participant's impression of product efficacy assessed using an efficacy questionnaire at 4, 8 and 12 weeks

7. Participant's impression of product usability assessed using a usability questionnaire at at 12 weeks

### Overall study start date

19/09/2019

### **Completion date**

15/03/2021

# Eligibility

### Key inclusion criteria

- 1. Korean female subjects aged between 30 and 60 years.
- 2. Dry skin on cheek (hydration value is below 48 arbirary units [AU] by Corneometer®)
- 3. Skin wrinkles of over grade 3 by DERMAPRO standard photograph
- 4. No chronic or acute disease, including skin disease
- 5. Signed informed consent for the study.
- 6. Cooperative and available for follow-up during the study period

### Participant type(s)

Healthy volunteer

### Age group

### Adult

**Lower age limit** 30 Years

**Upper age limit** 60 Years

Sex

Female

### Target number of participants

170 (85 in test group and 85 in placebo group)

### Total final enrolment

160

### Key exclusion criteria

1. Has consumed probiotics as dietary supplements, food or beverage products during the last 2 weeks

2. Pregnant, nursing or planning to become pregnant

3. Irritation or symptomatic allergy to food, including ingredients of cosmetic, medical and test products

4. Has taken oral or topical antibiotics during the previous 3 months

5. Has taken oral retinoid/steroid drugs or topical applications including steroids during previous 6 months

6. Has used functional cosmetics for improvement of skin wrinkle, hydration and elasticity within the last 3 months

7. Previous interventions at test site (e.g. skin decortications, Botox and other skin treatments)

8. Has participated in a previous study without an appropriate intervening period (3 months) between studies

9. Has disease which might affect the study (e.g. cardiovascular, kidney, liver, thyroid,

gastrointestinal disease, gout)

10. Any skin disease (e.g. atopic dermatitis) at test site

11. Any chronic disease (e.g. diabetes, asthma, high blood-pressure) or psychiatric disorder (e.g. depression, schizophrenia, alcoholism, drug addiction)

12. Take a medicine for treatment of obesity (e.g. antidepressants, anorectics), contraceptives, hormones or diuretics

13. Take excessive alcohol (over 30 g alcohol per day)

14. Sensitive or hypersensitive skin

15. Damaged skin in or around the test area, which includes sunburn, tattoos, scars or other disfiguration on the test area

16. Has an abnormal result in screening clinical chemical analysis by medical specialist 17. Has any problem which may interfere with the aim of the study as the judgment of the principal investigator

### Date of first enrolment

06/10/2020

# Date of final enrolment 18/12/2020

## Locations

**Countries of recruitment** Korea, South

#### Study participating centre DERMAPRO Ltd. 4F, 30, BangbaeJungang-ro Seocho-gu Seoul Korea, South 06684

### Sponsor information

**Organisation** Danisco Sweeteners Oy

**Sponsor details** Sokeritehtaantie 20 Kantvik Finland 02460 +358 10 431 2235 alvin.ibarra@iff.com

Sponsor type

Industry

# Funder(s)

**Funder type** Industry

**Funder Name** Danisco Sweeteners Oy

# **Results and Publications**

### Publication and dissemination plan

Planned publication in a high-impact peer-reviewed journal.

### Intention to publish date

31/12/2024

### Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are not expected to be made available due to the investigational product being a food supplement, thus there is no regulatory obligation to supply participant level data.

### IPD sharing plan summary

Not expected to be made available

### Study outputs

Output type	Details
<b>Basic results</b>	

**Date created** 05/12/2024

 Date added
 Pee

 09/01/2025
 No

Peer reviewed?

Patient-facing? No