# Effectiveness of low doses of yeast beta-glucan 1,3/1,6 in adults with moderate stress in fighting respiratory infections, reducing fatigue, boosting the immune system and improving digestive health

Submission date 11/07/2023	<b>Recruitment status</b> No longer recruiting	[X] Prospectively registered [X] Protocol
<b>Registration date</b> 04/08/2023	<b>Overall study status</b> Ongoing	<ul> <li>Statistical analysis plan</li> <li>Results</li> </ul>
Last Edited 27/01/2025	<b>Condition category</b> Other	<ul> <li>Individual participant data</li> <li>[X] Record updated in last year</li> </ul>

### Plain English summary of protocol

Background and study aims

Beta-glucans derived from fungi and yeast are known for their immune-modulating effects. It has been asserted that yeast beta-glucan significantly improves the function of the immune system by activating macrophages, one of the primary defences of the immune system. Its benefits have been extensively studied globally among athletes, stressed women, elderly people, healthy adults and children. Upper respiratory tract infections are the most common form of infection in every age group worldwide and have a significant negative impact on productivity, healthcare spending, and the economy. Based on published studies, the application of yeast beta glucans was proven as a possible therapeutic and preventive approach in managing and preventing recurrent respiratory tract infections. Therefore, this study aims to evaluate the effectiveness of yeast beta-glucan 1,3/1,6 on fatigue, respiratory infection, immune markers and gut health among adults with moderate stress.

### Who can participate?

People aged 18-59 years with moderate stress and common cold symptoms in the past 6 months

### What does the study involve?

Participants are randomly allocated to take either Yeast Beta Glucan (YBG) dose 1, dose 2 or dose 3 or a placebo (a substance that has no effect, used as a control). The study duration is 12 weeks. In this study, several assessments will be conducted at the screening level, baseline, the sixth week and the twelfth week. Other assessments that will be conducted in this study include blood profile, gut microbiome analysis and diet recall. A total of 20ml of blood will be collected to conduct the complete blood count including immune and oxidative stress biomarkers. In addition, several questionnaire forms will be used to assess the occurrence of upper respiratory tract infection symptoms, mood status and fatigue. In addition, an anthropometric evaluation will be carried out to assess the nutritional status of the participants.

What are the possible benefits and risks of participating? The benefits may not be direct. The supplement has the potential to strengthen and improve the immune system of the participants who receive this supplement. Participants may or may not experience side effects. Side effects may include changes in blood pressure, nausea, diarrhea and vomiting.

Where is the study run from? Mead Johnson Nutrition (Malaysia)

When is the study starting and how long is it expected to run for? January 2023 to September 2025

Who is funding the study? Mead Johnson Nutrition (Malaysia) Educational Grant

Who is the main contact? 1. Prof. Dr Suzana Shahar, suzana.shahar@ukm.edu.my 2. Nur Nadia Mohamad Habibullah, P126613@siswa.ukm.edu.my

# **Contact information**

**Type(s)** Principal Investigator

**Contact name** Prof Suzana Shahar

### Contact details

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# Additional identifiers

EudraCT/CTIS number Nil known

**IRAS number** 

**ClinicalTrials.gov number** Nil known

Secondary identifying numbers UKM PPI/111/8/JEP-2023-211

# Study information

### Scientific Title

Efficacy of low dosage yeast beta-glucan 1,3/1,6 on respiratory infection, fatigue, immune markers and gut health among adults with moderate stress

### **Study objectives**

1. The yeast beta glucan 1,3/1,6 supplementation is effective in improving upper respiratory tract infection symptoms and its related immune responses among adults with moderate stress in Klang Valley.

2. The yeast beta glucan 1,3/1,6 supplementation is effective in improving fatigue and psychological aspects among adults with moderate stress in Klang Valley.

3. The yeast beta glucan 1,3/1,6 supplementation is effective in improving gut health among adults with moderate stress in Klang Valley.

### Ethics approval required

Ethics approval required

### Ethics approval(s)

Approved 22/06/2023, Research Ethics Committee, The National University of Malaysia (Sekretariat Etika Penyelidikan Universiti Kebangsaan Malaysia, Tingkat 1, Blok Klinikal, Hospital Canselor Tuanku Mukhriz, Pusat Perubatan UKM, Jalan Yaacob Latif, Bandar Tun Razak 56000 Cheras, Kuala Lumpur, 56000, Malaysia; +60 (0)391455046; sepukm@ukm.edu.my), ref: UKM PPI /111/8/JEP-2023-211

### Study design

Multicenter interventional double-blinded randomized controlled trial

### Primary study design

Interventional

### Secondary study design

Randomised controlled trial

### Study setting(s)

Community, Hospital, University/medical school/dental school, Workplace

### Study type(s)

Efficacy

### Participant information sheet

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

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

Respiratory infections, fatigue, immune system, and digestive health of adults with moderate stress

### Interventions

Current interventions as of 30/01/2024:

This is a randomized, double-blind, placebo-controlled study designed to determine the efficacy of Yeast Beta Glucan 1,3/1,6 on respiratory infection, fatigue, immune markers and gut health among adults for 12 weeks.

This study requires participants to take either Yeast Beta Glucan (YBG) dose 1, YBG dose 2, YBG dose 3 or a placebo (a substance that has no therapeutic effect, used as a control). Once registered for the study, the eligible participants will undergo simple randomization to determine group allocation.

In this study, several assessments will be conducted at the screening level, baseline, the sixth week and the twelfth week. Other assessments that will be conducted in this study include blood profile, gut microbiome analysis and 3-day diet record. A total of 20 ml of blood will be collected to conduct the complete blood count including immune and oxidative stress biomarkers. In addition, several questionnaire forms will be used to assess the occurrence of upper respiratory tract infection (URTI) symptoms, mood status and fatigue. In addition, an anthropometric evaluation will be made to assess the nutritional status of the study participant.

### Previous interventions:

This is a randomized, double-blind, placebo-controlled study designed to determine the efficacy of Yeast Beta Glucan 1,3/1,6 on respiratory infection, fatigue, immune markers and gut health among adults for 3 months.

The supplement is in a 60 g sachet known as Provital Immuna Plus® is a nutritious milk that helps to support a healthy immune system. Finished products in the form of 60 mg sachet registration number 20-1-03444. The subjects will be taking either 204 mg and 100 mg of Yeast Beta

Glucan or placebo with 0 mg in the form of a 60 g sachet mixed with 160 ml of warm water until dissolved, two times daily (30 g each serving).

The placebo which will be used is a sensory-identical supplement in a sachet, composed of 60 mg maltodextrin.

Once registered for the study, the eligible participants will undergo simple randomization to determine group allocation. The study duration is 12 weeks. In this study, several assessments will be conducted at the screening level, the fourth week, the sixth week and the twelfth week. Other assessments that will be conducted in this study include blood profile, gut microbiome analysis and diet recall. A total of 20 ml of blood will be collected to conduct the complete blood count including immune and oxidative stress biomarkers. In addition, several questionnaire forms will be used to assess the occurrence of upper respiratory tract infection (URTI) symptoms, mood status and fatigue. In addition, an anthropometric evaluation will be made to assess the nutritional status of the study subjects.

### Intervention Type

Supplement

### Primary outcome measure

Current primary outcome measure as of 22/01/2024:

 Upper respiratory tract infection (URTI) episodes: occurrence, severity and duration measured using the Wisconsin Upper Respiratory Symptom Survey (WURSS)-21 at baseline, weeks 6 and 12.
 Fatigue measured using the Profile of Mood State (POMS) at baseline and week 6 and 12. 3. Blood immune markers: cytokines interleukin: [IL] IL-6, IL 8, IL-10, TNF-interferon [IFN]-γ, inflammation - CRP measured using blood samples at baseline and week 12.

4. Biomarkers and oxidative stress: oxidative stress (MDA, LPO), inflammatory biomarkers (iNOS, Cox2) measured at baseline and week 12.

5. Quality of life measured using Short Form-36 (SF-36) at baseline and weeks 4, 6 and 12.

Previous primary outcome measure:

1. Basic health profile (BMI, waist circumference, body fat percentage, complete blood count, fasting blood sugar, lipid profile, liver function test, renal function test) at baseline and week 12 2. Upper respiratory tract infection (URTI) episodes: occurrence, severity and duration measured using the Wisconsin Upper Respiratory Symptom Survey (WURSS)-21 at baseline and weeks 4, 6 and 12

3. Blood immune markers: cytokines interleukin: [IL] IL-6, IL 8, IL-10, TNF-interferon [IFN]-γ, inflammation - CRP measured using blood samples at baseline and week 12

### Secondary outcome measures

Current secondary outcome measures as of 22/01/2024:

1. The gut bacterial diversity (a diversity,  $\beta$  diversity, comparative analysis) and predicted functional analysis will be assessed at baseline and week 12

Previous secondary outcome measures:

1. Fatigue measured using the Profile of Mood State (POMS) at baseline and weeks 4, 6 and 12 2. Biomarkers and oxidative stress: oxidative stress (MDA, LPO), inflammatory biomarkers (iNOS, Cox2) measured at baseline and week 12

3. Quality of life measured using Short Form-36 (SF-36) at baseline and weeks 4, 6 and 12

4. Microbiome abundance and functional analysis ( $\alpha$  diversity,  $\beta$  diversity, comparative analysis) at baseline and week 12

Overall study start date 01/01/2023

**Completion date** 30/09/2025

# Eligibility

### Key inclusion criteria

Current inclusion criteria as of 16/01/2024:

- 1. 18-59 years old
- 2. In a stressful environment
- 3. Scoring 14-26 (moderate stress) on the Perceived Stress Scale (PSS-10)
- 4. Symptoms of common colds in the past 6 months
- 5. Willing to comply with protocol requirements

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### Participant type(s)

Healthy volunteer, Health professional, Employee

### Age group

Adult

### Lower age limit

18 Years

### Upper age limit

59 Years

### Sex

Both

Target number of participants

200

### Key exclusion criteria

1. Participant on immunosuppressant (i.e., steroids)

2. Use of immune-modifying medications or dietary supplements affecting the immune system (at least 2 weeks of washing period for inclusion)

3. Any autoimmune or health concerns

4. Uncontrolled comorbidities including diabetes, hypertension, cardiovascular disease, chronic kidney disease, chronic liver disease, active malignancy, endocrine disorder and cancer

5. Uncontrolled chronic respiratory illness including allergic rhinitis or asthma, presence of nasal ulcers/polyps

- 6. Body temperature >38C at study commencement
- 7. Pregnant/lactating female

8. On specific dietary requirements (diabetic diet/low salt diet/etc)

9. On prebiotics or probiotics supplementation (at least 2 weeks of washing period for inclusion) 10. On antibiotics washing period within 2 weeks for inclusion

- 11. Disabilities individuals- long-term physical, mental, intellectual and sensory impairment
- 12. Individuals without symptoms of common colds for the past 6 months
- 13. Allergy to yeast, oat, wheat, barley
- 14. Influenza vaccination for the past 1 year
- 15. Pneumococcal vaccine for the past 5 years
- 16. Participate in another research study

### Date of first enrolment

02/02/2024

# **Date of final enrolment** 30/11/2024

### Locations

### **Countries of recruitment** Malaysia

### Study participating centre H-care Wellness Center

61, Jalan Raja Muda Abdul Aziz, Chow Kit Kuala Lumpur Malaysia 50300

### Study participating centre

**Pusat Jantung & Paru-paru** Tingkat 8, Blok Klinikal Hospital Canselor Tuanku Muhriz Jalan Yaacob Latif, Bandar Tun Razak Kuala Lumpur Malaysia 56000

#### Study participating centre Faculty of Medicine UKM

Level 6, Preclinical Building Jalan Yaacob Latif, Bandar Tun Razak Kuala Lumpur Malaysia 56000

# Sponsor information

**Organisation** Mead Johnson Nutrition Malaysia

### Sponsor details

201, Jalan Tun Sambanthan, Brickfields Kuala Lumpur Malaysia 50470 +60 (0)14 668 3620 SoFie.Tan@reckitt.com **Sponsor type** Industry

Website https://www.reckitt.com/

# Funder(s)

Funder type Industry

Funder Name Mead Johnson Nutrition

Alternative Name(s) Mead Johnson

**Funding Body Type** Private sector organisation

**Funding Body Subtype** For-profit companies (industry)

**Location** United States of America

# **Results and Publications**

**Publication and dissemination plan** Planned publication in a high-impact peer-reviewed journal.

Intention to publish date 01/06/2025

### Individual participant data (IPD) sharing plan

The datasets generated and/or analysed during the current study will be published as a supplement to the results publication

### IPD sharing plan summary

Published as a supplement to the results publication

Study outputs					
Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Participant information sheet			25/07/2023	No	Yes

Protocol article

No