# Probiotic capsule trial

Submission date	Recruitment status  No longer recruiting	<ul><li>Prospectively registered</li></ul>		
18/04/2005		☐ Protocol		
Registration date 24/05/2005	Overall study status Completed	Statistical analysis plan		
		[X] Results		
<b>Last Edited</b> 23/05/2008	Condition category Nutritional, Metabolic, Endocrine	[] Individual participant data		

### Plain English summary of protocol

Not provided at time of registration

### Contact information

### Type(s)

Scientific

#### Contact name

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#### Contact details

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### Additional identifiers

Protocol serial number 121/6

## Study information

Scientific Title

### **Study objectives**

In non-diseased host the probiotic health claims could be assessed by improvement of some measurable laboratory indices of well-established physiological functions of host, e.g. markers of antioxidative defense system.

#### Ethics approval required

Old ethics approval format

### Ethics approval(s)

Not provided at time of registration

#### Study design

Randomised controlled trial

### Primary study design

Interventional

### Study type(s)

**Not Specified** 

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

Healthy subjects

#### **Interventions**

Probiotic capsule versus placebo capsule.

- 1. The study group members (8 males and 4 females) took three probiotic containing capsules (8.4 log colony forming unit [CFU] per capsule) two times daily (daily dose 9.2 log CFU) for three weeks.
- 2. The placebo group (7 males and 5 females) received identical capsules without the probiotic strain.

### Intervention Type

Drug

#### Phase

**Not Specified** 

### Drug/device/biological/vaccine name(s)

Probiotic

### Primary outcome(s)

Not provided at time of registration

### Key secondary outcome(s))

Not provided at time of registration

### Completion date

30/04/2004

## **Eligibility**

### Key inclusion criteria

Inclusion criteria included the wish to participate, no known health problems, no medical conditions requiring drug therapy, no other yoghurts or no special diets.

### Participant type(s)

**Patient** 

### Healthy volunteers allowed

No

### Age group

Adult

#### Sex

All

#### Key exclusion criteria

Subjects with a history of gastrointestinal tract (GIT) disease, food allergy and acute infection, use of any antimicrobial agent within the last month or use of any regular concomitant medication were excluded.

#### Date of first enrolment

01/01/2004

#### Date of final enrolment

30/04/2004

### Locations

#### Countries of recruitment

Estonia

## Study participating centre

Ravila 19

Tartu Estonia 50411

## Sponsor information

### Organisation

Estonian Science Foundation (Estonia)

## Funder(s)

### Funder type

Government

#### **Funder Name**

Estonian Science Foundation (base funding ref 0418 and 5327)

### Alternative Name(s)

Estonian Science Foundation, ETF

### **Funding Body Type**

Private sector organisation

### **Funding Body Subtype**

Trusts, charities, foundations (both public and private)

#### Location

Estonia

## **Results and Publications**

Individual participant data (IPD) sharing plan

### IPD sharing plan summary

Not provided at time of registration

### **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	04/08/2005		Yes	No