

Assessing the clinical benefit of food supplement, Genecol®, in subjects with joint pain at the lower, upper limbs or at the lumbar spine

Submission date 01/02/2010	Recruitment status No longer recruiting	<input type="checkbox"/> Prospectively registered
		<input type="checkbox"/> Protocol
Registration date 22/02/2010	Overall study status Completed	<input type="checkbox"/> Statistical analysis plan
		<input checked="" type="checkbox"/> Results
Last Edited 12/08/2013	Condition category Musculoskeletal Diseases	<input type="checkbox"/> Individual participant data

Plain English summary of protocol

Background and study aims

Joint pain is a major cause of disability in subjects aged over 50 years. Symptomatic pain relief can be obtained with painkillers such as paracetamol or non-steroidal anti-inflammatory drugs. These treatments, while generally safe when used at low doses and for a short time, can result in serious complications (gastrointestinal bleeding, kidney failure, coronary heart disease) when used for a long time or at higher doses and could obviously reduce the adherence to treatment. In subjects with joint symptoms, food supplements are often taken by patients with the aim to relieve pain and improve physical function. GENACOL®, a food supplement made of collagen hydrolysate, is a food supplement that claims to improve joint symptoms. The aim of this study is to assess whether taking this food supplement over 24 weeks increases the number of subjects with an improvement in joint pain and/or physical function symptoms.

Who can participate?

Men and women aged over 50 years with joint pain (hip, knee, elbow, shoulder, hand and/or lumbar spine).

What does the study involve?

Subjects were randomly allocated to receive either GENACOL® (three hard gel capsules per day) or a dummy/placebo (identical hard gel capsules, to be consumed in the same daily dosage). The target joint that was followed-up throughout the study was the most painful joint at the first visit. As the product tested is a food supplement, no accurate diagnosis of joint pain was performed.

What are the possible benefits and risks of participating?

A reduction of joint pain is expected in the GENACOL group. No specific risk is expected as this product is already marketed without significant adverse events.

Where is the study run from?
University of Liège (Belgium).

When is the study starting and how long is it expected to run for?
The study started in July 2009 and finished in January 2011.

Who is funding the study?
The study was financed by Nutraveris (France).

Who is the main contact?
Prof Jean-Yves Reginster
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Contact information

Type(s)
Scientific

Contact name
Prof Jean-Yves Reginster

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers
NUTRA-CH-01

Study information

Scientific Title
A randomised, double-blind, placebo-controlled phase IV trial to assess the clinical benefit of Genacol®, a food supplement made of a proprietary collagen hydrolysate (1200 mg/day), over a period of 24 weeks, in subjects with joint pain at the lower, upper limbs or at the lumbar spine

Study objectives

Genacol® is a food supplement made of a proprietary collagen hydrolysate already registered, as food supplement, in various countries, including Belgium, Canada, France, the United Kingdom, Spain, Italy, the United States, and others. The aim of this supplementation is to reduce pain and reduce functional disabilities in subjects with joint pain.

On 12/08/2013, the anticipated end date was changed from 01/07/2010 to 01/01/2011.

Ethics approval required

Old ethics approval format

Ethics approval(s)

The Ethics Committee of Liege CHU approved on 17/06/2009 (ref: B70720096310)

Study design

Randomised double blind 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 below to request a patient information sheet.

Health condition(s) or problem(s) studied

Joint pain

Interventions

Participants will be randomised to receive

1. Genacol®

Active ingredient: Collagene hydrolysate (400 mg/capsule; 1200 mg/day)

2. Placebo

Main ingredient: maltodextrin (400 mg/capsule; 1200 mg/day)

Participants will be required to take three hard gel capsules once daily before bed during the entire study period (24 weeks)

Intervention Type

Other

Phase

Phase IV

Primary outcome measure

Percentage of clinical responders between the active treatment and placebo groups

Secondary outcome measures

Outcomes will be evaluated at weeks 12 and 24

1. Pain rescue treatment consumption
2. Pain/function changes; assessed by
 - 2.1. Visual analogue scale (VAS)
 - 2.2. Lequesne index
 - 2.3. Disability of Arm Shoulder and Hand (DASH) score
 - 2.4. Functional Disability Scale for the Assessment of Low Back Pain (Echelle d'Incapacité Fonctionnelle pour l'Evaluation des Lombalgies [EIFEL]) questionnaire
3. Health-related quality of life changes; assessed with the SF-36
4. Utility value changes; assessed by EQ-5D
5. Tolerability and incidence of any adverse events
6. Global satisfaction of the treatment will be assessed by mean of a global questionnaire (also at week 4)

Overall study start date

01/07/2009

Completion date

01/01/2011

Eligibility

Key inclusion criteria

Men and women over 50 years with joint pain (hip, knee, elbow, shoulder, hand and lumbar spine) over 30 mm on a 0-100 mm visual analogue scale.

Participant type(s)

Patient

Age group

Senior

Sex

Both

Target number of participants

200

Key exclusion criteria

1. Unlikely to cooperate in the study
2. Poor compliance anticipated by the investigator
3. Participating in another trial at the same time or within the previous 1 month with active therapeutic intervention (except if the patient only performed the screening visit without taking the tested supplement)

Date of first enrolment

01/07/2009

Date of final enrolment

01/01/2011

Locations

Countries of recruitment

Belgium

Study participating centre

University of Liège

Liege

Belgium

4000

Sponsor information

Organisation

Nutraveris (France)

Sponsor details

Nutraveris

18 C, rue du Sabot

Ploufragan

France

22440

Sponsor type

Industry

ROR

<https://ror.org/05xc9cz73>

Funder(s)

Funder type

Industry

Funder Name

Nutraveris (France)

Results and Publications

Publication and dissemination plan

Not provided at time of registration

Intention to publish date

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	01/06/2012		Yes	No