

Clinical Trial Protocol

Iranian Registry of Clinical Trials

03 Jun 2026

Study of the effect of canola and olive oils consumption on anthropometric, inflammatory, glycemic and hormonal indices, lipid profiles, grade of fatty liver and mood disorder score in patients with polycystic ovary syndrome

Protocol summary

Study aim

Determination of the effects of canola and olive oil on anthropometric and inflammatory indices, glycemic and hormone indices, lipid profile, fatty liver grade and mood disturbance in patients with PCOS.

Design

The present study is a randomized double-blind clinical trial with control group on patients with PCOS.

Settings and conduct

90 samples were divided into three groups receiving 25 grams/day of Canola oil, 25 grams/day of olive oil and 25 grams/day of sunflower oil, using a randomized permuted block method. every groups recieved diet using adjusted ideal body weight. Macronutrient distribution in diet will be given in the form of 65-45% CHO, 10-15% Pr and 30-35% fat, and replace the calculated fat content with intervention and control oils. In this study, there are 3 visits for the patient at the beginning, week 5 and the end of the study. The intervention and control oils are given at a monthly rate with 25 grams modulus. This study was conducted in the form of double-blind that patients and physicians and investigators be unaware of the type of intervention that they were taking.

Participants/Inclusion and exclusion criteria

Inclusion criteria: -People with PCOS/18 to 45 years/overweight or obesity. -no use of any drug and / or surgical treatment -Not having any systemic disease and other endocrine disorders -Lack of Pregnancy and Breastfeeding Exclusion criteria: -Start taking or any dose changes in medications -Get pregnancy - Acceptance of less than 80% of the intervention

Intervention groups

In this study, two groups receiving 25 grams/day of canola oil and 25 grams/day of olive oil as intervention groups and one group receiving 25 grams/day of

sunflower oil as a control group.

Main outcome variables

Effects of canola and olive oil on anthropometric indices, inflammatory indices, glycemic and hormonal indices, lipid profile, lipid profile and mood disturbance in patients with PCOS

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20190407043193N1**

Registration date: **2019-06-30, 1398/04/09**

Registration timing: **retrospective**

Last update: **2019-06-30, 1398/04/09**

Update count: **0**

Registration date

2019-06-30, 1398/04/09

Registrant information

Name

Maryam Yahay

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 31 3662 4711

Email address

maryam.yahay@gmail.com

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2018-10-22, 1397/07/30
Expected recruitment end date
2019-03-15, 1397/12/24
Actual recruitment start date
2018-10-22, 1397/07/30
Actual recruitment end date
2019-03-15, 1397/12/24
Trial completion date
2019-03-17, 1397/12/26

Scientific title

Study of the effect of canola and olive oils consumption on anthropometric, inflammatory, glycemic and hormonal indices, lipid profiles, grade of fatty liver and mood disorder score in patients with polycystic ovary syndrome

Public title

effect of canola and olive oils consumption in polycystic ovary syndrome

Purpose

Treatment

Inclusion/Exclusion criteria

Inclusion criteria:

The definitive diagnosis of PCOS based on diagnostic criteria (Rotterdam3) by the physician The absence of other endocrine disorders or the conditions that lead to some or most of the clinical manifestations and disorders associated with PCOS (such as congenital or non-classic adrenal hyperplasia, Cushing's syndrome, androgens secreting tumors, drugs induced hyperandrogenism, idiopathic hyperandrogenism, hirsutism Idiopathic, thyroid dysfunction, hyperprolactinemia, pregnancy, lactation and menopause) Non-use of any drug and / or surgical treatment for the clinical symptoms and disorders associated with PCOS other than OCP (such as spironolactone, finasteride, isotretinoin, letrozole, clomiphene, gonadotropins, metformin, cyproterone, rosiglitazone, pioglitazone, ovarian laparoscopic surgery, and Auxiliary reproductive technology) Absence of any severe or significant systemic disease requiring treatment such as any cancer; digestive, liver or endocrine disorders (such as celiac disease, Crohn's disease, ulcerative colitis), diabetes mellitus, hyperparathyroidism, hypercalcemia, or hyperphosphatemia); cardiovascular disorders (e.g. Uncontrolled hypertension or history of myocardial infarction); kidney disorders (such as renal failure, nephrotic syndrome); blood coagulation disorders (eg thalassemia, hemophilia); neurological disorders (such as epilepsy) or reproductive disorders associated with PCOS Ask a doctor or a patient Not taking any of these items: tobacco; alcohol; anti-estrogens (such as tamoxifen and raloxifene); oral or injectable corticosteroids (such as prednisone, prednisolone, dexamethasone, triamcinolone, hydrocortisone or betamethasone); lack of omega-3 supplementation Chains and long chains); effective drugs for insulin resistance such as metformin and sitagliptin Non-consumption of any canola and olive oil in the last 6 months as the main consumer oil Lack of any allergy, intolerance or harmful drug reaction to the supplementation of the studied oils Being in the age range of 18-45 years (people are not better off at the age

of growth and menopause) Being in the BMI range above 25 and less than 40 Ability to understand the goals of the study and provide informed written consent The desire to participate in the study Weight constant over the past 6 months (self report) Lack of Pregnancy and Breastfeeding

Exclusion criteria:

The onset of a dose or any change in dosage in the above drugs or the change in the type or dose of OCP consumed during the study period Getting pregnant during the study period Incidence of severe side effects or signs of poisoning with supplements used during the study period Failure to adhere to the study protocol The acceptance of less than 80% of the intervention (consumption of less than 80% of the total intervention oil that should be consumed during the 10-week intervention period will be considered as a low admission)

Age

From **18 years** old to **45 years** old

Gender

Female

Phase

2-3

Groups that have been masked

- Participant
- Care provider
- Investigator
- Outcome assessor

Sample size

Target sample size: **90**

Actual sample size reached: **90**

Randomization (investigator's opinion)

Randomized

Randomization description

The samples were classified using randomized block splitting and divided into three groups using random numbers table.

Blinding (investigator's opinion)

Double blinded

Blinding description

In this study, the patients were randomly divided into three groups who were informed about entering the study and were not aware of the type of intervention received. On the other hand, the researcher who does all the information and measurements is also unaware of the type of intervention that each patient receives. A collaborator physician in the study was also blinded to this study.

Placebo

Not used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics Committee of Isfahan University of Medical Sciences

Street address

Isfahan University of Medical Sciences. Hazar Jarib Street

City

isfahan

Province

Isfahan

Postal code

8174673461

Approval date

2019-11-17, 1398/08/26

Ethics committee reference number

IR.MUI.RESEARCH.REC.1397.315

Health conditions studied

1

Description of health condition studied

Polycystic ovary syndrome

ICD-10 code

E28.2

ICD-10 code description

Polycystic ovarian syndrome

Primary outcomes

1

Description

lipid profile

Timepoint

At the beginning of the study, 5 weeks after the intervention and the end of the study

Method of measurement

Spectrophotometric method

2

Description

Anthropometric Indicators

Timepoint

At the beginning of the study, 5 weeks after the intervention and the end of the study

Method of measurement

Measurements with biometric impedance analysis (BIA) and meters

3

Description

fatty liver grade

Timepoint

At the beginning of the study, 5 weeks after the

intervention and the end of the study

Method of measurement

sonography

4

Description

mood disorder score

Timepoint

At the beginning of the study, 5 weeks after the intervention and the end of the study

Method of measurement

DASS-21 questionnaire

5

Description

Inflammatory Indicators

Timepoint

At the beginning of the study, 5 weeks after the intervention and the end of the study

Method of measurement

ELISA

Secondary outcomes

1

Description

Clinical signs including hirsutism and the interval between menstruation

Timepoint

At the beginning of the study, 5 weeks after the intervention and the end of the study

Method of measurement

MFGS questionnaire And the question of the patient

2

Description

blood pressure

Timepoint

At the beginning of the study, 5 weeks after the intervention and the end of the study

Method of measurement

Mercury barometric

Intervention groups

1

Description

Intervention group: 25 grams of olive oil

Category

Treatment - Other

2

Description

Intervention group: 25 grams of canola oil

Category

Treatment - Other

3

Description

Control group: 25 grams of sunflower oil

Category

Treatment - Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Martyrs' clinic

Full name of responsible person

akbar samadi

Street address

Martyrs' clinic. Sepahanshahr. Nezam Street.

City

isfahan

Province

Isfahan

Postal code

8173763738

Phone

+98 31 3651 0020

Email

info@shohadaclinic.ir

Web page address

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Esfahan University of Medical Sciences

Full name of responsible person

shaghaiegh haghjoo

Street address

Isfahan University of Medical Sciences. Hazar Jarib Street

City

isfahan

Province

Isfahan

Postal code

8174673461

Phone

+98 31 3668 0048

Email

mui@gmail.com

Grant name

Thesis

Grant code / Reference number

Is the source of funding the same sponsor organization/entity?

Yes

Title of funding source

Esfahan University of Medical Sciences

Proportion provided by this source

60

Public or private sector

Public

Domestic or foreign origin

Domestic

Category of foreign source of funding

empty

Country of origin

Type of organization providing the funding

Academic

Person responsible for general inquiries

Contact

Name of organization / entity

Esfahan University of Medical Sciences

Full name of responsible person

Maryam Yahay

Position

Phd student

Latest degree

Medical doctor

Other areas of specialty/work

Nutrition

Street address

Hezarjrib ave, Isfahan university of medical science. faculty of nutrition.

City

Esfahan

Province

Isfahan

Postal code

8156685691

Phone

+98 31 3662 4711

Fax

Email

maryam.yahay@gmail.com

Person responsible for scientific inquiries

Contact

Name of organization / entity

Esfahan University of Medical Sciences

Full name of responsible person

Maryam Yahay

Position

Phd student

Latest degree

Medical doctor

Other areas of specialty/work

Nutrition

Street address

Hezarjrib ave, Isfahan university of medical science. faculty of nutrition.

City

Esfahan

Province

Isfahan

Postal code

8156685691

Phone

+98 31 3662 4711

Fax

Email

maryam.yahay@gmail.com

Person responsible for updating data

Contact

Name of organization / entity

Esfahan University of Medical Sciences

Full name of responsible person

Maryam Yahay

Position

Phd student

Latest degree

Medical doctor

Other areas of specialty/work

Nutrition

Street address

Hezarjrib ave, Isfahan university of medical science.
faculty of nutrition.

City

Esfahan

Province

Isfahan

Postal code

8156685691

Phone

+98 31 3662 4711

Fax

Email

maryam.yahay@gmail.com

Sharing plan

Deidentified Individual Participant Data Set (IPD)

Yes - There is a plan to make this available

Study Protocol

Yes - There is a plan to make this available

Statistical Analysis Plan

Yes - There is a plan to make this available

Informed Consent Form

Yes - There is a plan to make this available

Clinical Study Report

Yes - There is a plan to make this available

Analytic Code

Undecided - It is not yet known if there will be a plan to make this available

Data Dictionary

Undecided - It is not yet known if there will be a plan to make this available

Title and more details about the data/document

Proposals and consent form and published article from the study

When the data will become available and for how long

Getting Started October 1399

To whom data/document is available

Academic Institutions

Under which criteria data/document could be used

To give students access to their dissertations

From where data/document is obtainable

You can call 09134118836 for help

What processes are involved for a request to access data/document

After doing the research and publishing the paper, it will provide you with the documentation.

Comments