

Improving the long-term immune response to the flu vaccine in older people

Submission date 09/08/2024	Recruitment status No longer recruiting	<input type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
Registration date 30/10/2024	Overall study status Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
Last Edited 28/02/2025	Condition category Other	<input type="checkbox"/> Individual participant data <input checked="" type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

Influenza is a great threat to public health and most deaths occur in older people. Effective vaccination protecting older people from flu is a global health challenge. Long-lasting immunity following flu vaccination is difficult to achieve, only 52% of older people have protective antibodies after vaccination. The immune system becomes less efficient with age and immune responses to vaccines reduce. We want to confirm our exciting preliminary results showing that a dietary supplement, made from wheatgerm, can restore and improve levels of polyamine intake in older people. Older people have low levels due to reduced dietary intake. Polyamines can be absorbed from natural food types, e.g., wheatgerm and pomegranates. However, a large volume of food needs to be consumed to increase the level of polyamine dietary intake. A double-blinded, randomised study we conducted showed that the supplement is safe when given after a COVID vaccine and that immune biomarkers increase in older people, with a statistical significance compared to placebo. Our primary outcome will determine if this holds true when the supplement is given after a licensed flu vaccine. We will determine the feasibility of recruitment to the trial, retention of participants, tolerability of supplements, adverse reactions and the side-effect profile of the supplement when taken after the flu vaccine. Our secondary outcome will be exploratory, measuring immune biomarkers: antibodies, T/B cells, after the flu vaccine. We will compare the effects of the supplement to that of a placebo supplement. Blood samples will be analysed at Cardiff University and University of Oxford. This study will justify a Phase 1 clinical trial, where the supplement is administered with the flu vaccine to study the longevity of the immune response and measure clinical outcomes. This research develops health knowledge relating to flu prevention and enables people to manage their dietary intake at home with an oral food supplement.

Who can participate?

People who are 65 years and over are able to participate.

What does the study involve?

Participants taking part in the study will attend 4 research appointments over 37 weeks, at baseline, week 2, week 13 and week 37.

1. Following informed consent, at the baseline appointment participants will be asked questions about their health and these recorded in a form, have their pulse, blood pressure, oxygen levels, temperature and weight measured.

At the baseline visit they will receive a licensed flu vaccine. They will donate 60 mls of venous blood at baseline appointment for biochemistry and haematology profiles and immunogenicity assays.

They will be given a placebo or polyamine supplement to take home and instructions on how to take this for 2 weeks and asked to fill in a diary of missed doses and side effects.

2. At the week 2 visit, participants will be asked questions about their health and these recorded in a form, have their pulse, blood pressure, oxygen levels, temperature and weight measured.

They will donate 60 mls of venous blood at baseline appointment for biochemistry and haematology profiles and immunogenicity assays. Missed doses or side effects will be recorded by the research team. They will be given a placebo or polyamine supplement to take home and instructions on how to take this for 11 weeks and asked to fill in a diary of missed doses and side effects.

3. At the week 13 visit, participants will be asked questions about their health and these recorded in a form, have their pulse, blood pressure, oxygen levels, temperature and weight measured. They will donate 50 mls of venous blood for immunogenicity assays. Missed doses or side effects will be recorded by the research team. They will be given a placebo or polyamine supplement to take home and instructions on how to take this for 11 weeks and asked to fill in a diary of missed doses and side effects.

4. At the week 37 visit, participants will be asked questions about their health and these recorded in a form, have their pulse, blood pressure, oxygen levels, temperature and weight measured. They will donate 50 mls of venous blood for immunogenicity assays. This is the end of the study.

Participants will be able to claim expenses at the end of the study and will be informed of the study results.

What are the possible benefits and risks of participating?

We cannot guarantee there are any benefits to taking part in the study.

Where is the study run from?

Cardiff University (UK)

When is the study starting and how long is it expected to run for?

May 2024 to November 2025

Who is funding the study?

Health and Care Research Wales (UK)

Who is the main contact?

Dr Lucy Jones, jonesl147@cardiff.ac.uk

Contact information

Type(s)

Public, Scientific, Principal Investigator

Contact name

Dr Lucy Jones

ORCID ID

<http://orcid.org/0000-0002-3872-4376>

Contact details

Department of Microbiology, 6th Floor, University hospital of Wales
Cardiff
United Kingdom
CF14 4XN
02920742394
JonesL147@cardiff.ac.uk

Additional identifiers

EudraCT/CTIS number

Nil known

IRAS number

314310

ClinicalTrials.gov number

Nil known

Secondary identifying numbers

SPON1966-23, CPMS 64340

Study information

Scientific Title

IRFLUVA: Improving the long-term immune response to the flu vaccine in older people. A double-blinded, randomised controlled feasibility trial of the effects of a polyamine rich food supplement on flu vaccine antibody levels and immune cell responses, compared to a placebo supplement, in people aged 65 years and over.

Acronym

IRFLUVA

Study objectives

Is a wheat-germ extract supplement safe after a flu vaccine and can it improve the amplitude and longevity of antibody and cellular immune response to the flu vaccine in people aged over 65 years?

Ethics approval required

Ethics approval required

Ethics approval(s)

Approved 07/05/2024, Black Country HRA Research Ethics Committee (2 Redman Place, Stratford, London, E20 1JQ, United Kingdom; +44 2071048129; blackcountry.rec@hra.nhs.uk), ref: 24/WM/0065

Study design

Interventional non-CTIMP single-centre double-blind randomized controlled study

Primary study design

Interventional

Secondary study design

Randomised controlled trial

Study setting(s)

Hospital, University/medical school/dental school

Study type(s)

Safety, Efficacy

Participant information sheet

<https://blogs.cardiff.ac.uk/irfluva-study/volunteers-wanted-for-a-healthy-immune-ageing-study-2/>

Health condition(s) or problem(s) studied

Healthy ageing. Immune ageing

Interventions

Arm 1: Polyamine supplement – 6mg supplement taken orally for 13 weeks after a licensed flu vaccine. Follow-up for 37 weeks in total.

Arm 2: Placebo – 6mg of a rice flour placebo taken orally for 13 weeks after a licensed flu vaccine. Follow-up for 37 weeks in total.

Randomisation using REDCAP randomisation tool. The trial is double-blinded with stratified randomisation by biological sex.

Intervention Type

Supplement

Primary outcome measure

Feasibility measures:

1. Data from the IRFLUVA study case report forms will be used to measure study recruitment rates and retention rates.
2. Data from the IRFLUVA study case report forms will be used to measure study withdrawals and the reason for withdrawal.
3. Data from the IRFLUVA study case report forms will be used to measure the number, severity and nature of adverse reactions for all participants at baseline, 2 weeks, 13 weeks and 37 weeks.
4. Data from the IRFLUVA study case report form will be used to measure the number, severity and nature of adverse events for all participants at baseline, 2 weeks, 13 weeks and 37 weeks.
5. Data from the IRFLUVA study case report form will be used to measure Compliance with polyamine supplement and placebo supplements from baseline to week 13.
6. Results from biochemistry and haematology profile testing at baseline, 2 weeks, 13 weeks and

37 weeks.

7. The number of protocol deviations will be measured at the end of the study.

Secondary outcome measures

Measured for up to 37 weeks:

1. T cell Elispots to measure responses to flu
2. B cell Elispots to measure B cell responses to flu
3. IgG ELISA to measure antibodies to flu
4. Plaque assays to study neutralizing antibodies
5. Cytokine assays to measure cytokine responses

Overall study start date

07/05/2024

Completion date

30/11/2025

Eligibility

Key inclusion criteria

1. 65 years of age or over
2. Capacity to provide written informed consent
3. Eligible for a seasonal flu vaccine
4. Willing to receive a licensed flu vaccine as part of the study protocol on visit 1 of the protocol

Participant type(s)

Healthy volunteer

Age group

Senior

Lower age limit

65 Years

Upper age limit

101 Years

Sex

Both

Target number of participants

80

Key exclusion criteria

1. Acutely unwell
2. Had a flu vaccine in the 6 months prior to recruitment
3. Known to be immunosuppressed
4. History of influenza in the 6 months prior to recruitment
5. Cannot provide informed written consent
6. Not willing to receive a licensed flu vaccine as part of the study protocol on visit 1 of the

protocol

7. Use systemic steroids for more than one week e.g., prednisolone >0.5mg/kg/day in the three months prior to first study intervention
8. Chronic administration (≥ 14 days in total) of immunosuppressants or other immune modifying drugs in the 3 months prior to first study intervention
9. Receipt of blood, blood products and/or plasma derivatives or any immunoglobulin preparation in the three months prior to first study intervention
10. Diagnosed with diabetes
11. Diagnosed with allergy to, or constituent parts of, SpermidineLife supplements or who have gluten intolerance
12. Already taking Spermidine supplements at the time of recruitment or for 6 months prior to recruitment to the study
13. In custody
14. Currently participating in other interventional trials
15. Do not live in Wales

Date of first enrolment

30/09/2024

Date of final enrolment

30/10/2024

Locations

Countries of recruitment

United Kingdom

Wales

Study participating centre

University Hospital of Wales

Heath Park

Cardiff

United Kingdom

CF14 4XW

Sponsor information

Organisation

Cardiff University

Sponsor details

Park Place

Cardiff

Wales

United Kingdom

CF10 3AT
+44 2920742394
FalconerHE@cardiff.ac.uk

Sponsor type

University/education

Website

<http://www.cardiff.ac.uk/>

ROR

<https://ror.org/03kk7td41>

Funder(s)

Funder type

Government

Funder Name

Health and Care Research Wales

Alternative Name(s)

Health & Care Research Wales, Ymchwil Iechyd a Gofal Cymru, Health Care Research Wales, HCRW

Funding Body Type

Government organisation

Funding Body Subtype

Local government

Location

United Kingdom

Results and Publications

Publication and dissemination plan

Results from the study will be disseminated to the participants in the form of a letter. Public engagement events will be used to disseminate results. Online platforms and Communication channels will be used to share results. Results will be presented at scientific conferences and published in peer-reviewed journals.

Intention to publish date

30/11/2026

Individual participant data (IPD) sharing plan

Anonymised datasets generated and analysed during the study will be available upon request from Dr Lucy Jones JonesL147@cardiff.ac.uk following publication.

IPD sharing plan summary

Available on request