Investigating the effects of recommended - but unproven - therapies for anaphylaxis in a human histamine challenge model

Submission date 02/08/2024	Recruitment status Recruiting	[X] Prospectively registered [] Protocol
Registration date	Overall study status	
05/08/2024	Ongoing	☐ Results
Last Edited	Condition category	☐ Individual participant data
05/08/2024	Other	Record updated in last year

Plain English summary of protocol

Background and study aims

Anaphylaxis is a life-threatening allergic reaction that happens very quickly. Epinephrine and other interventions are recommended for the treatment of anaphylaxis. However, there is no evidence for effectiveness from randomized controlled trials. The aim of this trial is to compare the treatments recommended in anaphylaxis guidelines in a randomised controlled trial in healthy volunteers challenged with histamine.

Who can participate?

Healthy volunteers aged 18 years and over

What does the study involve?

Healthy volunteers will undergo a histamine challenge to lower blood pressure in order to test interventions that may raise blood pressure (injections of adrenaline, infusion of fluid and elevation of legs).

What are the possible benefits and risks of participating?

As histamine has rapidly reversible actions the risks are limited mainly to blood sampling and side effects such as flushing and headache. There will be no direct benefit for the participants other than information form the performed health tests.

Where is the study run from? Medical University of Vienna (Austria)

When is the study starting and how long is it expected to run for? March 2022 to January 2030

Who is funding the study? Investigator initiated funding not yet secured

Contact information

Type(s)

Public, Scientific, Principal investigator

Contact name

Prof Bernd Jilma

Contact details

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Additional identifiers

Clinical Trials Information System (CTIS)

2022-003591-16

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

Nil known

Study information

Scientific Title

Histamine infusion in healthy volunteers to study changes in biomarkers and effects of therapeutic interventions – a pilot study

Acronym

EPI/HIS

Study objectives

Histamine challenge can be used as a surrogate to examine therapies to treat anaphylaxis

Ethics approval required

Ethics approval required

Ethics approval(s)

approved 21/03/2022, Medical University of Vienna (Borschkegasse, Vienna, 1090, Austria; +43 (0)140400; ethik-kom@meduniwien.ac.at), ref: IRB00002503

Study design

Partly randomized controlled trial

Primary study design

Interventional

Study type(s)

Treatment, Efficacy

Health condition(s) or problem(s) studied

Shock

Interventions

Recommended interventions for anaphylaxis treatment; this is part of the adaptive trial design and may change. Additional arms may be added in the platform trial approach

- 1. Elevation of legs (10 min)
- 2. Volume of crystalloids 10 ml/kg over 10 min
- 3. Control injection of placebo (same volume as adrenaline)
- 4. Adrenaline doses as recommended in current anaphylaxis guidelines

Intervention Type

Mixed

Primary outcome(s)

Mean arterial blood pressure measured with an automated device over 15 minutes during histamine infusion

Key secondary outcome(s))

- 1. Diastolic and systolic blood pressure with an automated device measured over 15 minutes during histamine infusion
- 2. Pharmacokinetics of adrenaline measured with qualified mass spectrometry assays over 60 minutes

Completion date

01/01/2030

Eligibility

Key inclusion criteria

Healthy volunteers

Participant type(s)

Healthy volunteer

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

All

Key exclusion criteria

- 1. Non-healthy
- 2. Pregnant
- 3. Breastfeeding

Date of first enrolment

15/08/2024

Date of final enrolment

01/01/2030

Locations

Countries of recruitment

Austria

Study participating centre Medical University of Vienna

Dept of Clinical Pharmacology Währingergerürtel 18-20 Wien Austria 1090

Sponsor information

Organisation

Medical University of Vienna

ROR

https://ror.org/05n3x4p02

Funder(s)

Funder type

Other

Funder Name

Investigator initiated funding not yet secured

Results and Publications

Individual participant data (IPD) sharing plan

Due to data protection issues, only anonymized data will be shared with qualified researchers. Prof. Bernd Jilma (bernd.jilma@meduniwien.ac.at) should be contacted for access to datasets, anonymized raw data of the primary and secondary endpoints will be shared with qualified investigators once the related manuscripts have been published, and the request has been cleared by the institution's data protection unit.

IPD sharing plan summary

Available on request, Other

Study outputs

Output type Details Date created Date added Peer reviewed? Patient-facing?

Participant information sheet Participant information sheet 11/11/2025 11/11/2025 No Yes