

Eczema monitoring online via questionnaires

Submission date 21/06/2021	Recruitment status No longer recruiting	<input checked="" type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
Registration date 29/06/2021	Overall study status Completed	<input type="checkbox"/> Statistical analysis plan <input checked="" type="checkbox"/> Results
Last Edited 11/07/2024	Condition category Skin and Connective Tissue Diseases	<input type="checkbox"/> Individual participant data

Plain English summary of protocol

Background and study aims

Eczema is a chronic inflammatory skin condition that affects children and adults. It is characterised by periods of remission and relapse, indicating the fluctuating nature of the disease. Online questionnaires completed by participants are often used in eczema clinical trials to capture participant's views about their eczema. This allows researchers to measure the effectiveness of different treatments. It is possible that completing questionnaires regularly (known as monitoring) can change the way that people manage their eczema and so result in improved eczema severity. This might be due to the completion of the questionnaires triggering participants to think about their eczema more or to apply treatments more often. This is a concern within clinical trials because completing regular questionnaires may make it harder to identify changes in eczema resulting from the treatments being tested. This study will assess whether completing questionnaires regularly changes participants' behaviour and affects eczema severity. The overall aim of this study is to inform the design of future eczema clinical trials about the effect of monitoring.

Who can participate?

Patients aged 1 year or older with eczema

What does the study involve?

Participants will be split into two groups: the intervention group will be asked to complete questionnaires weekly for 8 weeks, and the control group will be asked to complete questionnaires at the beginning and end of the study only. The online questionnaire takes about 10 minutes to complete. This will allow the researchers to evaluate the effect of regular monitoring on eczema severity.

What are the possible benefits and risks of participating?

There will be no direct benefit to participants, but their participation will help to improve future eczema research. Participating in this study will allow participants to track their eczema symptoms at home, which they may find useful and interesting. There are no anticipated risks to participants from taking part. Participants will be able to use their normal eczema treatment throughout this research study.

Where is the study run from?

Centre of Evidence-Based Dermatology at the University of Nottingham (UK)

When is the study starting and how long is it expected to run for?
January 2021 to April 2022

Who is funding the study?
University of Nottingham (UK)

Who is the main contact?
Arabella Baker
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Contact information

Type(s)
Scientific

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Additional identifiers

Clinical Trials Information System (CTIS)
Nil known

ClinicalTrials.gov (NCT)
Nil known

Protocol serial number
EMO 120421

Study information

Scientific Title
Evaluation of the effect of symptom monitoring with patient-reported outcome measures on clinical outcomes in eczema: an online, parallel-group randomised controlled trial (EMO trial)

Acronym
EMO

Study objectives

Weekly symptom monitoring will enhance self-management of eczema, which in turn will improve adherence to standard treatment use, which will lead to improved eczema severity at 8 weeks.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved 23/04/2021, Faculty of Medicine & Health Science Research Ethics Committee, University of Nottingham (Faculty Hub, Room E41, E Floor, Medical School, Queen's Medical Centre Campus, Nottingham University Hospitals, Nottingham, NG7 2UH, UK; +44 (0) 115 74 85060; FMHS-ResearchEthics@nottingham.ac.uk), ref: 239-0421

Study design

Online parallel-group randomized controlled clinical trial

Primary study design

Interventional

Study type(s)

Other

Health condition(s) or problem(s) studied

Eczema

Interventions

This is a methodological trial and there will be no treatment intervention involved. The intervention will be the online Patient Oriented Eczema Measure (POEM) questionnaire. Eligible participants will be randomised online and assigned to either the intervention group or to the control group using the REDCap software. The randomisation schedule will be based on computer-generated codes using random permuted blocks of varying size, stratified by age and baseline disease severity. The PhD student will deal with participant queries, thus she will have access to group allocation, but the other trial team members and the trial statistician will remain blinded. Participants in the intervention group (weekly group) will be asked to complete the POEM questionnaire weekly for 8 weeks. The control group will not receive the intervention.

Intervention Type

Other

Primary outcome(s)

Eczema severity measured by the self-reported or proxy (parent/carer reported) POEM questionnaire score at baseline and 8 weeks

Key secondary outcome(s)

1. Adherence to eczema treatment use, assessed in two ways:
 - 1.1. Eczema treatment use assessed by the emollient and topical corticosteroid use questionnaire over the last week from baseline to 8 weeks

1.2. Eczema treatment use assessed by the overall eczema treatment use questionnaire 2 months prior to joining the study and at the end of the study at 8 weeks

2. Missing data: the proportion of fully completed questionnaires at 8 weeks

Completion date

03/04/2022

Eligibility

Key inclusion criteria

1. Self-report or parent/carer report of eczema diagnosis by a healthcare professional
2. Person aged 1 year or older
3. Able and willing to provide informed consent
4. If under 16 years old, a parent/carer needs to provide informed consent on behalf of the child
5. Able to read and understand written English
6. Have access to the internet and to an internet-enabled device
7. POEM score of 3 or above at eligibility screening

Healthy volunteers allowed

No

Age group

Mixed

Sex

All

Total final enrolment

296

Key exclusion criteria

1. Unable/unwilling to provide informed consent
2. Currently taking part in another eczema clinical trial

Date of first enrolment

14/09/2021

Date of final enrolment

16/01/2022

Locations

Countries of recruitment

United Kingdom

England

Afghanistan

Åland Islands

Albania

Algeria

American Samoa

Andorra

Angola

Anguilla

Antarctica

Antigua and Barbuda

Argentina

Armenia

Aruba

Australia

Austria

Azerbaijan

Bahamas

Bahrain

Bangladesh

Barbados

Belarus

Belgium

Belize

Benin

Bermuda

Bhutan

Bolivia

Bonaire Saint Eustatius and Saba

Bosnia and Herzegovina

Botswana

Bouvet Island

Brazil

British Indian Ocean Territory

Brunei Darussalam

Bulgaria

Burkina Faso

Burundi

Cabo Verde

Cambodia

Cameroon

Canada

Cayman Islands

Central African Republic

Chad

Chile

China

Christmas Island

Cocos (Keeling) Islands

Colombia

Comoros

Congo

Congo, Democratic Republic

Cook Islands

Costa Rica

Croatia
Cuba
Curaçao
Cyprus
Czech Republic
Côte d'Ivoire
Denmark
Djibouti
Dominica
Dominican Republic
Ecuador
Egypt
El Salvador
Equatorial Guinea
Eritrea
Estonia
Eswatini
Ethiopia
Falkland Islands
Faroe Islands
Fiji
Finland
France
French Guiana
French Polynesia
French Southern Territories

Gabon
Gambia
Georgia
Germany
Ghana
Gibraltar
Greece
Greenland
Grenada
Guadeloupe
Guam
Guatemala
Guernsey
Guinea
Guinea-Bissau
Guyana
Haiti
Heard Island and McDonald Islands
Holy See (Vatican City State)
Honduras
Hong Kong
Hungary
Iceland
India
Indonesia
Iran

Iraq
Ireland
Isle of Man
Israel
Italy
Jamaica
Japan
Jersey
Jordan
Kazakhstan
Kenya
Kiribati
Korea, North
Korea, South
Kosovo
Kuwait
Kyrgyzstan
Lao People's Democratic Republic
Latvia
Lebanon
Lesotho
Liberia
Libya
Liechtenstein
Lithuania
Luxembourg

Macao

Madagascar

Malawi

Malaysia

Maldives

Mali

Malta

Marshall Islands

Martinique

Mauritania

Mauritius

Mayotte

Mexico

Micronesia, Federated States of

Moldova

Monaco

Mongolia

Montenegro

Montserrat

Morocco

Mozambique

Myanmar

Namibia

Nauru

Nepal

Netherlands

New Caledonia

New Zealand

Nicaragua

Niger

Nigeria

Niue

Norfolk Island

North Macedonia

Northern Mariana Islands

Norway

Oman

Pakistan

Palau

Palestine, State of

Panama

Papua New Guinea

Paraguay

Peru

Philippines

Pitcairn

Poland

Portugal

Puerto Rico

Qatar

Romania

Russian Federation

Rwanda

Réunion

Saint Barthélemy

Saint Helena, Ascension and Tristan da Cunha

Saint Kitts and Nevis

Saint Lucia

Saint Martin (French part)

Saint Pierre and Miquelon

Saint Vincent and the Grenadines

Samoa

San Marino

Sao Tome and Principe

Saudi Arabia

Senegal

Serbia

Seychelles

Sierra Leone

Singapore

Sint Maarten (Dutch part)

Slovakia

Slovenia

Solomon Islands

Somalia

South Africa

South Georgia and the South Sandwich Islands

South Sudan

Spain
Sri Lanka
Sudan
Suriname
Svalbard and Jan Mayen
Sweden
Switzerland
Syria
Taiwan
Tajikistan
Tanzania
Thailand
Timor-Leste
Togo
Tokelau
Tonga
Trinidad and Tobago
Tunisia
Turkmenistan
Turks and Caicos Islands
Tuvalu
Türkiye
Uganda
Ukraine
United Arab Emirates
United States Minor Outlying Islands

United States of America

Uruguay

Uzbekistan

Vanuatu

Venezuela

Viet Nam

Virgin Islands, British

Virgin Islands, U.S.

Wallis and Futuna

Western Sahara

Yemen

Zambia

Zimbabwe

Study participating centre

University of Nottingham

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Sponsor information

Organisation

University of Nottingham

ROR

<https://ror.org/01ee9ar58>

Funder(s)

Funder type

University/education

Funder Name

University of Nottingham

Alternative Name(s)

The University of Nottingham

Funding Body Type

Private sector organisation

Funding Body Subtype

Universities (academic only)

Location

United Kingdom

Results and Publications

Individual participant data (IPD) sharing plan

All data generated or analysed during this study will be included in the subsequent results publication.

IPD sharing plan summary

Other

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	17/05/2023	17/05/2023	Yes	No
Other publications	Effect on recruitment of social media strategy	27/10/2022	31/10/2022	Yes	No
Participant information sheet	version v1.0	09/04/2021	08/07/2021	No	Yes
Protocol file	version 1.0	08/04/2021	18/08/2022	No	No