

Low-dose radon hyperthermia therapy in atopic dermatitis

Submission date 24/06/2021	Recruitment status No longer recruiting	<input 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 type="checkbox"/> Results
Last Edited 27/08/2024	Condition category Skin and Connective Tissue Diseases	<input type="checkbox"/> Individual participant data
		<input type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

Atopic dermatitis (AD) is an inflammatory skin disease, mainly characterized by pruritus, skin dryness, and eczema. For some patients, conventional therapy methods are insufficient. These patients might benefit from Low-dose Radon Hyperthermia (LDRnHT) therapy. This is a well-recognized treatment method for inflammatory diseases in various compartments of the human body. Clinical studies demonstrated that LDRnHT- therapy can reduce pain, enhance functionality and positively shift crucial blood parameters. Many positive single reports from AD patients support this theory, but there are no clinical studies so far. The purpose of this pilot study is to evaluate whether AD patients can evidently benefit from LDRnHT.

Who can participate?

Patients aged 18 -70 years with chronic, moderate to severe AD

What does the study involve?

All study participants have to pass an initial examination (T0) at the Department of Dermatology and Allergology at the University Hospital Salzburg, Austria. Patients enrolled are randomised into a control and an intervention group. Participants attend a two-week cure stay in Bad Gastein. The intervention group receives eight sessions of LDRnHT in the Gastein Healing Gallery (average radon activity ≈ 44 kBq/m³; ambient temperature 37-41.5° C; air humidity $\geq 70\%$). The control group receives sauna treatments with the same ambient temperature and humidity but without radon. Short term modifications of skin condition and specific blood parameters are assessed, as well as questionnaires for skin condition and quality of life (Qol), comparing the initial situation (T0) to immediate post-radon-therapy (T1).

Long-term effects are documented in follow-up examinations at three, six and nine months after T1 (T2/T3/T4, respectively). At every examination, skin condition is assessed by a dermatologist using the SCORing Atopic Dermatitis rating tool, and blood samples and questionnaires are collected. Patients are asked to fill out the Patient Oriented SCORAD questionnaire using a computer or smartphone application, and the SKINDEX-29 tool to measure of the effect of the skin disease on quality of life. The EQ-5D-5L and VAS questionnaires are used to assess Quality of Life (Qol). At T0 and T2. If the condition of a participant becomes worse, the dermatologist is authorized to assign a different therapy and end a patient's participation in the study. In this pilot study, molecular parameters shall be investigated which reflect the therapeutic

effects of LDRnHT in AD patients. The results from this pilot study shall build a basis for the development of further hypotheses and larger studies investigating LDRnHT as a therapy option for AD.

What are the possible benefits and risks of participating?

Patients receive a cost-free cure stay in Bad Gastein. Improvement of skin condition might be achieved through LDRnHT.

Possible worsening of skin condition might occur due to elevated air humidity and temperature (ambient temperature 37-41.5°C; air humidity \geq 70%) during treatments.

Where is the study run from?

The study is run from the Gastein Research Institute (Paracelsus Medical University in Salzburg) which is part of the Center for Physiology, Pathophysiology and Biophysics (Salzburg and Nuremberg, Austria and Germany), in cooperation with the Gastein Healing Gallery in Bockstein, Austria.

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

August 2016 to December 2020

Who is funding the study?

Cure stays and treatment funded by:

Gasteiner Kur-, Reha und Heilstollen Betriebsges.m.b.H. (Austria)

Other funding:

Gastein Research Institute, Center for Physiology, Pathophysiology and Biophysics, Salzburg and Nuremberg - Salzburg, Paracelsus Medical University Salzburg (Austria)

Gemeinnützige Salzburger Landeskliniken Betriebsgesellschaft mbH (Austria)

Who is the main contact?

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Additional identifiers**Protocol serial number**

E2126

Study information**Scientific Title**

Study for the effectivity of low-dose radon hyperthermia therapy in the Gastein healing gallery on atopic dermatitis

Study objectives

This pilot study outcome shall provide information for further generation of hypotheses

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved 25/01/2017, Ethikkommission für das Bundesland Salzburg (Sebastian-Stief-Gasse 2 5020 Salzburg, Austria), +43-(0)662-8042-2375, ethikkommission@salzburg.gv.at), ref: E.-Nr. 2126

Study design

Single-centre interventional open randomized explorative prognostic longitudinal pilot study

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Effectivity of low-dose radon hyperthermia therapy on atopic dermatitis

Interventions

Patients with Atopic Dermatitis (AD) are recruited via the department of dermatology of the Salzburger Landeskliniken (Prim. Dr. Johann Bauer, SALK).

Patients are randomised to control (sauna treatment) and interventional groups (Low-dose Radon Hyperthermia Therapy (LDRnHT) in the Gastein Healing Gallery).

The randomisation process was carried out, regardless of any specific patient characteristics, via permuted block randomisation (block sizes: 7, 6, and 4).

Blood samples, questionnaires and survey of drug consumption were collected for timepoints before, directly after cure stay and three, six, and nine months after end of the cure stay.

Intervention Type

Procedure/Surgery

Primary outcome(s)

SCORAD (SCORing Atopic Dermatitis) questionnaire for specific assessment of skin condition in AD at before, directly after cure stay and 3, 6, and 9 months after the end of the cure stay.

Key secondary outcome(s)

Measured before, directly after cure stay, and three, six, and nine months after the end of the cure stay:

1. Quality of life (EQ-5D-5L)
2. Skin condition (SKINDEX-29)
3. Analysis of blood samples (MDC/CCL22, CTACK/CCL27, TARC/CCL17, IgE, IL-4, IL-13)

Completion date

08/12/2020

Eligibility

Key inclusion criteria

1. Atopic Dermatitis, from chronic moderate to severe disease state. (mild: SCORAD <25, moderate SCORAD 25-50, severe SCORAD >50)
2. Patient SCORAD value must be between 25 and 50 for inclusion
3. AD must be diagnosed and verified by a dermatologist
4. Age range 18 - 70 years
5. Signed Informed Consent must be obtained from each patient
6. Patients must be able to fill out a questionnaire by themselves

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Upper age limit

70 years

Sex

All

Total final enrolment

34

Key exclusion criteria

1. Pregnancy or breast feeding period
2. Incompatibility of patients to treatment methods
3. Erosions, ulcers
4. Viral or bacterial superinfections
5. Severe internal diseases
6. Consumption of potential photosensitizers
7. Concomitant or past malignant skin tumors
8. Radon therapy or similar radiation (UV/light) treatment within the past year
9. Consumption of oral immunosuppressive drugs (within 6 months before entering the study)
10. Claustrophobia

Date of first enrolment

25/01/2017

Date of final enrolment

06/07/2020

Locations

Countries of recruitment

Austria

Germany

Study participating centre

Gasteiner Kur-, Reha- und Heilstollen Betriebsges.m.b.H.

Heilstollenstraße 19

Böckstein

Bad Gastein

Austria

5645

Study participating centre

Gastein Research Institute, Center for Physiology, Pathophysiology and Biophysics, Salzburg and Nuremberg - Salzburg, Paracelsus Medical University Salzburg

Strubergasse 22

Salzburg

Austria

5020

Study participating centre

Gemeinnützige Salzburger Landeskliniken Betriebsgesellschaft mbH

Müllner Hauptstraße 48

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Sponsor information

Organisation

Salzburger Landeskliniken

ROR

<https://ror.org/0500kmp11>

Organisation

Salzburger Landeskliniken

ROR

<https://ror.org/0500kmp11>

Organisation

Paracelsus Medizinische Privatuniversität

ROR

<https://ror.org/022zhm372>

Funder(s)

Funder type

University/education

Funder Name

Paracelsus Medizinische Privatuniversität

Alternative Name(s)

Paracelsus Medical University, PMU

Funding Body Type

Private sector organisation

Funding Body Subtype

Universities (academic only)

Location

Austria

Funder Name

Gasteiner Kur-, Reha und Heilstollen Betriebsges.m.b.H.

Funder Name

Gemeinnützige Salzburger Landeskliniken Betriebsgesellschaft mbH

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?
Participant information sheet	version v4	12/06/2019	08/07/2021	No	Yes
Protocol file	In German	31/12/2017	10/08/2022	No	No