# ChILD-EU database and observational study

<b>Submission date</b> 07/11/2013	<b>Recruitment status</b> No longer recruiting	[X] Prospectively registered [_] Protocol
<b>Registration date</b> 19/12/2013	<b>Overall study status</b> Completed	<ul> <li>[] Statistical analysis plan</li> <li>[X] Results</li> </ul>
Last Edited 28/09/2020	<b>Condition category</b> Respiratory	Individual participant data

### Plain English summary of protocol

Background and study aims

Childhood Interstitial Lung Diseases (ChILD) are a group of rare diseases of the lung: most conditions have a poor outcome. There are too few cases in each country to enable adequate research. The ChILD-EU project, funded by the European Commission, is bringing together clinicians and chILD cases from across Europe. The study will gather information into a Europewide database and also enable outcomes to be studied.

Who can participate?

Infants and children coming to hospital with suspected interstitial lung disease

What does the study involve?

Information is collected on each patient at diagnosis, who are then observed over the first year following diagnosis (at 1, 2, 3, 6 and 12 months). At the time of diagnosis all patients in the database have their diagnosis and treatment reviewed by an expert team to ensure diagnostic validity. Measurements recorded typically are those routinely monitored during normal clinic visits. Parents and older children are also asked to fill in questionnaires at the start of the study and again after 3, 6 and 12 months. To enable genetic investigation, blood samples are collected from each child and their parents.

What are the possible benefits and risks of participating? There are no direct benefits to the parents or children taking part in this study. However, the information from this study will show which approaches to treatment give better outcomes.

Where is the study run from? The study is run from hospitals across Europe

When is the study starting and how long is it expected to run for? January 2014 to June 2016

Who is funding the study? European Commission Directorate-General for Research and Innovation, FP7-Health-2012-Innovation-1 Who is the main contact? Dr Steve Cunningham Steve.cunningham@ed.ac.uk

**Study website** http://www.childeu.net

## **Contact information**

**Type(s)** Scientific

**Contact name** Dr Steve Cunningham

**ORCID ID** http://orcid.org/0000-0001-7342-251X

## Contact details

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## Type(s)

Public

**Contact name** Ms Morag MacLean

ORCID ID http://orcid.org/0000-0002-4037-0247

## **Contact details**

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## Additional identifiers

EudraCT/CTIS number

**IRAS number** 

ClinicalTrials.gov number

Secondary identifying numbers N/A

## Study information

## Scientific Title

Orphans Unite: ChILD better together EUropean management platform for childhood interstitial lung diseases

## Acronym

ChILD-EU

## Study objectives

There are limited studies bringing together children with interstitial lung disease and no studies assessing the response to standardised interventions in Childhood Interstitial Lung Diseases (ChILD). The paucity of cases in each centre and the lack of an evidence-based treatment approach requires a structured observation of current practice to inform future research directions. The aim is to capture interventions and outcomes in well-characterised patients with suspected and proven ChILD. Such information will provide data on outcome in relation to standard interventions and support further research directions.

Studies in France, Germany, Italy and Turkey will collect similar information to add to the UK data in the database.

## Ethics approval required

Old ethics approval format

Ethics approval(s) South-East Scotland REC2, 08/11/2013, ref: 13/SS/0195

**Study design** Observational cohort multi-centre study

**Primary study design** Observational

**Secondary study design** Cohort study

**Study setting(s)** Hospital

**Study type(s)** Screening

### Participant information sheet

Not available in web format, please use the contact details to request a patient information sheet

## Health condition(s) or problem(s) studied

Childhood interstitial lung disease (chILD)

## Interventions

Observations will start from time of presentation at hospital during which the diagnosis is made and participants will continue in the trial for 12 months. Participants will be given the usual treatment for ChILD and data will be collected at seven time points. The data collected will include respiratory measurements, treatments, images of scans and histology samples and patient-reported outcome questionnaires. At study entry blood samples for genetic analysis will be collected from the participant and the participant's parents. Previously diagnosed cases of chILD will only enter the database and biobank study and so will only capture data at study entry and for peer review after 1 year.

Intervention Type

Other

**Phase** Not Applicable

### Primary outcome measure

For the database and biobank study - collate detailed information on clinical cases of possible ChILD on a central database and biobank.

For the observational study - describe outcomes at 1, 2, 3, 6 and 12 months in infants and children with ChILD.

Outcomes measured will be:

1. Death

- 2. Survival on artificial ventilatory support (invasive or non-invasive)
- 3. Survival in supplemental oxygen
- 4. Survival breathing room air
- 5. Quality of life (QoL)

## Secondary outcome measures

For the database and biobank study:

1. To review each case by an experienced international interdisciplinary peer review team to provide diagnostic oversight and feedback

- 2. To provide annual updates of diagnosis and outcome in a feedback loop via peer review
- 3. To store for future research, blood samples for genetic analysis of cases and parents
- 4. To support paediatricians and families caring for children with ChILD

For the observational study:

To describe variance in outcome at 1, 2, 3, 6 and 12 months in infants and children with ChILD according to:

- 1. Diagnosis and presentation
- 1.1. Diagnosis (peer review)
- 1.2. Diagnostic certainty (peer review)
- 1.3. Computed tomography (CT) score by component radiologist (peer review)

- 1.4. Blood oxygen saturation (SpO2) at rest in room air at presentation
- 1.5. SpO2 asleep in room air at presentation (nadir)
- 1.6. Respiratory rate (RR) (z score) at rest in air at presentation
- 1.7. Heart rate (HR) (z score) in air at presentation
- 1.8. Blood pressure at rest for 5 minutes at presentation
- 1.9. Weight (z-score) at presentation

1.10. Leland Fan 5 point severity score (nil, symptoms, SpO2 <90% air asleep, SpO2 at rest, pulmonary hypertension).

2. Time to treatment and improvement

2.1. Time from onset of symptoms/signs of ChILD to first treatment

2.2. Time from onset of symptoms/signs of ChILD to diagnosis (local clinical)

2.3. Time from onset of symptoms/signs of ChILD to normoxia whilst awake (SpO2 ≥94% breathing room air at rest)

2.4. Time from onset of symptoms/signs of ChILD to respiratory rate in normal range for age (Fleming, Thompson et al. 2011)

2.5. Time from onset of first treatment to reduction in RR by 10%

2.6. Time from onset of first treatment to reduction in HR by 20%

2.7. Time from onset of symptoms/signs of ChILD to normoxia whilst asleep (SpO2 ≥94% breathing room air at rest)

2.8. Time from onset of symptoms/signs of ChILD to weight appropriate for age/height without use of calorie supplementation

2.9. Time from onset of treatment to improvement in weight by 10%

3. Treatments

3.1. Steroids: use of steroids, dose, route and frequency of steroid use, time from first presentation to initiation of steroids, number of concomitant ChILD treatments at time of starting steroids

3.2. Hydroxychloroquine: use of hydroxychloroquine, dose and frequency of

hydroxychloroquine, time from first presentation to initiation of hydroxychloroquine, number of concomitant ChILD treatments at time of starting hydroxychloroquine

3.3. Azithromycin: use of azithromycin, dose and frequency of azithromycin, time from first presentation to initiation of azithromycin, number of concomitant ChILD treatments at time of starting azithromycin

4. Concomitant medicines

5. Follow-up review

5.1. SpO2 in room air measured 4 weeks after commencing initial treatment

5.2. RR at rest measured 4 weeks after commencing initial treatment

5.3. Heart rate at rest measured 4 weeks after commencing initial treatment

6. Quality of Life score - PEDS QL Generic Core Scales at 0 and 12 months

7. Questionnaire for health care utilisation and costs

7.1. Utilisation of inpatient and outpatient care to calculate direct costs gathered at 0, 3, 6 and 12 months

7.2. Loss of productivity of parents and children to calculate indirect costs gathered at 0, 3, 6 and 12 months

### Overall study start date

01/12/2013

## **Completion date**

30/11/2016

## Eligibility

### Key inclusion criteria

Infants and children presenting to hospital with clinician-suspected interstitial lung disease or at least three of the following four criteria present:

1. Respiratory symptoms for at least 14 days

1.1. Cough

- 1.2. Rapid and/or difficult breathing
- 1.3. Exercise intolerance
- 2. Respiratory signs
- 2.1. Tachypnea
- 2.2. Adventitious sounds
- 2.3. Retractions
- 2.4. Digital clubbing
- 2.5. Failure to thrive
- 2.6. Respiratory failure
- 3. Hypoxemia
- 4. Diffuse abnormalities on a chest radiograph or computerised tomography (CT) scan

Participant type(s)

Patient

Age group

Child

Sex

Both

**Target number of participants** 32

### Total final enrolment

127

### Key exclusion criteria

A participant would be excluded from the database if ineligible to participate in the ChILD-EU Minimal Dataset observation and follow-up study.

Exclusion criteria are common causes of diffuse lung disease, including but not exclusively: 1. Cystic fibrosis

- 2. Respiratory distress syndrome
- 3. Bronchopulmonary dysplasia
- 4. Acute infection (viral or bacterial)
- 5. Inherited or acquired immune deficiency

### Date of first enrolment

01/04/2014

Date of final enrolment 30/11/2016

## Locations

## Countries of recruitment

England

Scotland

United Kingdom

Wales

**Study participating centre Royal Hospital for Sick Children** Edinburgh United Kingdom EH9 1LF

#### **Study participating centre John Radcliffe Hospital** Headley Way Oxford United Kingdom OX3 9DU

#### **Study participating centre Royal Liverpool Children's Hospital** Alder Hey Eaton Road Liverpool United Kingdom L12 2AP

#### **Study participating centre King's College Hospital** Denmark Hill London United Kingdom SE5 9RS

**Study participating centre Leeds General Infirmary** Great George Street Leeds United Kingdom LS1 3EX

**Study participating centre RCPCH Nottingham Children's Hospital** QMC Derby Road Nottingham United Kingdom NG7 2UH

**Study participating centre Royal Aberdeen Children's Hospital** Cornhill Road Aberdeen United Kingdom AB25 2ZG

**Study participating centre Bristol Royal Hospital for Children** Upper Maudlin Street Bristol United Kingdom BS2 8BJ

**Study participating centre Royal Brompton Hospital** Sydney Street London United Kingdom SW3 6NP

**Study participating centre The Royal Hospital for Children** 1345 Govan Road Govan United Kingdom G51 4TF

#### **Study participating centre The Royal Victoria Infirmary** Queen Victoria Road

Newcastle upon Tyne United Kingdom NE1 4LP

#### **Study participating centre Royal Manchester Children's Hospital** Oxford Road Manchester United Kingdom M13 9WL

### Study participating centre

**Sheffield Children's Hospital** Western Bank Sheffield United Kingdom S10 2TH

## Study participating centre Birmingham Children's Hospital

Steelhouse Lane Birmingham United Kingdom B4 6NH

#### **Study participating centre Great Ormond Street Hospital for Children** Great Ormond Street London United Kingdom WC1N 3JH

### Study participating centre

Southampton General Hospital Tremona Road Southampton United Kingdom SO16 6YD **Study participating centre Noah's Ark Children's Hospital for Wales** University Hospital of Wales Heath Park Cardiff United Kingdom CF14 4XW

**Study participating centre Royal London Hospital** Whitechapel Road Whitechapel London United Kingdom E1 1BB

## Sponsor information

**Organisation** Academic and Clinical Centre Office for Research and Development (ACCORD) (UK)

### Sponsor details

University of Edinburgh & NHS Lothian The Queens Medical Research Institute 47 Little France Crescent Edinburgh United Kingdom EH16 4TJ

**Sponsor type** Research organisation

Website http://www.accord.ed.ac.uk/

ROR https://ror.org/01x6s1m65

## Funder(s)

Funder type

### Funder Name

European Commission Directorate-General for Research and Innovation, FP7-Health-2012-Innovation-1, Funding ref nr 305653

## **Results and Publications**

### Publication and dissemination plan

Planned publication in a high-impact peer-reviewed journal within one year after the end of the trial.

### Intention to publish date

30/11/2017

### Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be stored in a nonpublically available repository – the Child-EU registry which is administered by the Kids Lung Register Foundation. Access to anonymised datasets should be requested from the Kids Lung Registry Foundation. Contact ChILD-EU.register@med.uni-muenchen.de and Prof Matthias Griese (Matthias.Griese@med.uni-muenchen.de) for further information.

### IPD sharing plan summary

Stored in repository

### Study outputs

Output type	<b>Details</b> results	Date created	Date added	Peer reviewed?	Patient-facing?
Results article		01/02/2020	28/09/2020	Yes	No
HRA research summary			28/06/2023	No	No