

# European Health Examination Survey

<https://www.neurodegenerationresearch.eu/survey/european-health-examination-survey/>

## Title of the cohort

European Health Examination Survey

## Acronym for cohort

LUX-EHES

## Name of Principal Investigator

Title mme

First name Marie-Lise

Last name LAIR

## Address of institution where award is held

Institution

Street Address 1A-B rue Thomas Edison

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## Country

Luxembourg

## Website

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## Contact email

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## Funding source

Health Ministry

Research Ministry

## 1. The cohort includes, or expects to include, incidence of the following conditions

- Neurodegenerative disease in general

## When studies on the above condition(s) are expected to become possible

2011 – 2015

## 2a. Stated aim of the cohort

to study public health determinants

## **2b. Features distinguishing this cohort from other population cohorts**

we ask blood sample for biobanking

### **3a. i) Number of publications that involve use of cohort to date**

0

### **3a. ii) Up to three examples of studies to date (PI, Institution, Title of Study)**

### **3b. Publication list/link to where data or publications are accessible (if available)**

### **3c. Information (i.e. research findings) expected to be gained from the population cohort**

epidemiological informations public health informations impact of environmental conditions on health and some diseases

### **4a. Study criteria: age range of participants at recruitment**

Age in years from: more 18 years

To ('until death' if applicable): 65 years

### **4b. Study criteria: inclusion criteria**

criteria from european health examination survey will be used. People will be randomly selected from the national register.

### **4c. Study criteria: exclusion criteria**

institutionalised people

## **5. Size of the cohort (i.e. number of participants enrolled)**

5,001 – 10,000 participants

### **6a. Measures used to characterise participants**

age, gender, comorbidities, socio economic conditions, socio educative conditions, environmental conditions, lifestyle (tobacco, nutrition, alcohol, physical activity), medications, diseases, health status, family diseases, use of healthcare, ethnical informations, work conditions, vaccines, anthropometric measurements, blood pressure, blood analysis, visual capacity, auditiv capacity, cognitiv status

### **6b. Additional measures for participants with a clinical disorder**

### **6c. Are there defined primary and secondary endpoints (e.g. defined health parameters)**

No

## **7. Study design**

Cross sectional survey

## **8. Cases matched by**

- Age
- Sex
- Co-morbidities
- Cognitive function
- Physical ability

**9a. Does the study include a specialised subset of control participants**

No

**9b. If yes, description of specialised subset of control participants**

**10a. i) Data collection start date**

01-01-2012

**10a. ii) Data collection end date**

31-12-2013

**10a iii) Data collection for this study is**

- At the planning stage

**10b. Plans to continue the cohort study beyond the current projected end date**

- Yes – intend to apply for funding

**11. Data collected**

- Only through the study

**12. System in place to enable re-contact with patients for future studies**

Yes (participants have given permission to be re-contacted via the PIs to ask if they would participate in further studies)

**13a. Format and availability of data stored in a database**

Yes/No % available

Data summarised in database yes

Database is web-based yes

Database on spreadsheet

Database is on paper

Other (specify)

**Language used:**

french, german, english, portuges

**13b. Format and availability of data held as individual records**

**Language used:**

**14a. Are data available to other groups**

Yes

**14b. Access policy/mechanisms for access if data are available to other groups**

- Apply to PI or co-ordinator at resource
- Access through collaboration with PI only
- Access Committee mechanism
- Local/ regional access
- National access
- International access
- Access restricted to peer-reviewed work
- Applicant needs to provide separate external ethics approval

**15. Data sharing policy specified as a condition of use**

Data made publicly available after a specified time point

**16a. Are tissues/samples/DNA available to other groups**

Yes

**16b. i) Description of available tissues/samples/DNA**

- Living donors: blood
- Living donors: blood derivatives
- Living donors: DNA

**16b. ii) Form available tissues/samples/DNA are supplied in**

**16b. iii) Is the access policy/mechanism for obtaining samples the same as that for obtaining data**

Yes

**17. Is information on biological characteristics available to other groups**

No