

MEMENTO: Determining Factors and The Progression of The Onset of Alzheimer's Disease and Cognitive Impairment - Cohort of Patients with A Cognitive Complaint

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| General | |
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| Identification | |
| Detailed name | Cohort of Patients with A Cognitive Complaint |
| Sign or acronym | MEMENTO: Determining Factors and The Progression of The Onset of Alzheimer's Disease and Cognitive Impairment |
| CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation | CPP: 2010-A01394-35 (15/12 2010); AFSSAPS: B101404-30 (6/12/2010). |
| General Aspects | |
| Medical area | Geriatrics Neurology |
| Health determinants | Social and psychosocial factors |
| Keywords | loss of independence, cognitive decline, quality of life |
| Scientific investigator(s) (Contact) | |
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| Organization | Institut de la santé et de la recherche médicale - |
| Name of the director | Dufouil |

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| Organization | Institut de la santé et de la recherche médicale |
| Collaborations | |
| Funding | |
| Funding status | Mixed |
| Details | Alzheimer's Foundation Plan (Fondation Plan Alzheimer), PHRC |
| Governance of the database | |
| Sponsor(s) or organisation(s) responsible | CHU de Bordeaux |
| Organisation status | Public |
| Additional contact | |
| Main features | |
| Type of database | |
| Type of database | Study databases |
| Study databases (details) | Cohort study |
| Database recruitment is carried out by an intermediary | A selection of health institutions and services |
| Database recruitment is carried out as part of an interventional study | No |
| Additional information regarding sample selection. | Patients recruited from memory resource and research centres (CM2R) in France; regional expertise and clinical research centres for cognitive impairment. |
| Database objective | |
| Main objective | To study the onset of early signs (cognitive complaints, psychometric test performances lower than normal) that may indicate Alzheimer's disease or a related illness. |
| Inclusion criteria | - Male or female; |

- Adult;
- With a cognitive complaint.

Population type

Age

Adulthood (19 to 24 years)
 Adulthood (25 to 44 years)
 Adulthood (45 to 64 years)
 Elderly (65 to 79 years)
 Great age (80 years and more)

Population covered Sick population

Gender

Male
 Woman

Geography area National

Detail of the geography area France

Data collection

Dates

Date of first collection (YYYY or MM/YYYY) 2011

Size of the database

Size of the database (number of individuals) [1000-10 000[individuals

Details of the number of individuals 1,928 (enrolment on 21/02/14).

Data

Database activity Current data collection

Type of data collected

Clinical data
 Paraclinical data
 Biological data

Clinical data (detail) Direct physical measures

Details of collected clinical data ----

Paraclinical data (detail) Neuropsychological tests, positron emission tomoscintigraphy and MRI.

Biological data (detail) Lumbar puncture.

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| Presence of a biobank | Yes |
| Contents of biobank | Fluids (saliva, urine, amniotic fluid, ?) |
| Details of biobank content | Cerebrospinal fluid. |
| Health parameters studied | Health event/mortality Quality of life/health perception |
| Procedures | |
| Data collection method | Clinical, biological, psychological, sociological and brain imaging data will be collected throughout the study. |
| Classifications used | ---- |
| Quality procedure(s) used | ---- |
| Participant monitoring | Yes |
| Monitoring procedures | Monitoring by convocation of the participant |
| Details on monitoring of participants | ---- |
| Links to administrative sources | No |
| Promotion and access | |
| Promotion | |
| Link to the document | Memento.pdf |
| Access | |
| Terms of data access (charter for data provision, format of data, availability delay) | The cohort is a translational research platform open to sub-studies and ancillary studies submitted by any researcher in good faith. One main data set will be available 6 months after each wave of study monitoring and data cleaning. Please contact the scientist in charge of the study for access to more specific data. |
| Access to aggregated data | Free access |
| Access to individual data | Access on specific project only |