

MFV - Cross-sectional study on Migrations-Family-Ageing in French overseas departments

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General

Identification

Detailed name Cross-sectional study on Migrations-Family-Ageing in French overseas departments

Sign or acronym MFV

CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation CNIL

General Aspects

Medical area General practice

Health determinants Geography
Healthcare system and access to health care services
Lifestyle and behavior
Social and psychosocial factors

Keywords French overseas department, Demography, Intergenerational, Language practices, Religious practices, Migrations, Discrimination, Family, Ageing, Health

Scientific investigator(s) (Contact)

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Organization INED

Collaborations

Funding

Funding status Public

Details INED

Governance of the database

Sponsor(s) or organisation(s) responsible INED

Organisation status Public

Sponsor(s) or organisation(s) responsible Insee

Organisation status Public

Presence of scientific or steering committees Yes

Additional contact

Main features

Type of database

Type of database Study databases

Study databases (details) Not-repeated cross-sectional studies (except case control studies)

Database recruitment is carried out by an intermediary A population file

Database recruitment is carried out as part of an interventional study No

Additional information regarding sample selection. The survey was conducted among men and women between the ages of 18 and 79, in a representative sample of 4,000 households in each department, i.e. a total of 16,000 for the four French overseas departments.

Database objective

Main objective The MFV survey was carried out in a coordinated way in all the French overseas departments and focuses on the changes in family life, the sources and obstacles of social itineraries, the impact of migration and the evolution of intergenerational

solidarity

Inclusion criteria	Three main categories of key player were asked: * Native people of the department * Returning migrants: native people of the department who have had an emigration experience of more than six consecutive months and who were resettled there at the time of the survey * Immigrants: the non-native people of the department who were settled there at the time of the survey, whether French or foreign
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Population type

Age	Adulthood (19 to 24 years) Adulthood (25 to 44 years) Adulthood (45 to 64 years) Elderly (65 to 79 years)
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Population covered	General population
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Pathology

Gender	Male Woman
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Geography area	Local
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French regions covered by the database	Guadeloupe Guyane Martinique La Réunion
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Detail of the geography area	Guadeloupe (GP) French Guiana (GY) Martinique (MQ) Réunion (RE)
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Data collection

Dates

Date of first collection (YYYY or MM/YYYY)	2009
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Date of last collection (YYYY or MM/YYYY)	2010
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Size of the database

Size of the database (number of individuals)	[10 000-20 000] individuals
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Details of the number of individuals	15 770
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Data

Database activity Data collection completed

Type of data collected Declarative data

Declarative data (detail) Face to face interview

Details of collected declarative data THE EXTENDED FAMILY MODULE AND FAMILY SOLIDARITES (Members of the Ego family currently migrating, Solidarity and caregivers, State of health and access to health care) ; THE FAMILY MODULE (The children of Ego and his spouse, The present union and past unions of Ego, Mastery and the future of fertility); THE MODULE ON THE PROFESSIONAL AND MIGRATORY TRAJECTORIES OF NATIVE PEOPLE AND IMMIGRANTS (Social and professional trajectories, The professional and migratory trajectories of the native people of the department, The migratory and professional trajectories of people born outside the department, Experiences of rejection or discrimination in the department); THE CULTURAL RESOURCES MODULE (Citizen life, Languages, Religion)

Presence of a biobank No

Health parameters studied Health event/morbidity
Health care consumption and services
Quality of life/health perception

Care consumption (detail) Medicines consumption

Procedures

Data collection method CAPI questionnaire

Participant monitoring No

Followed pathology

Links to administrative sources No

Promotion and access

Promotion

Access

Terms of data access (charter for data provision, format of Contact the scientist in charge

data, availability delay)

Access to aggregated data

Access on specific project only

Access to individual data

Access on specific project only