COLOMBES - Louis Mourier Cohort: Severe and Morbid Obesity: Assessment, Evaluation and Follow-Up

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General	
Identification	
Detailed name	Louis Mourier Cohort: Severe and Morbid Obesity: Assessment, Evaluation and Follow-Up
Sign or acronym	COLOMBES
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	CNIL: 11/04/2006
General Aspects	
Medical area	Endocrinology and metabolism
Health determinants	Nutrition
Scientific investigator(s) (Contact)	
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Unit	SERVICE DE CHIRURGIE, HÔPITAL LOUIS MOURIER AP-HP
Organization	APHP
Collaborations	
Participation in projects, networks and consortia	Yes
Funding	
Funding status	Public
Details	AP-HP, direction de la recherche clinique - AP-HP, unité de recherche clinique Paris nord - AP-HP, hôpital Louis Mourier
Governance of the database	
	AP-HP
Sponsor(s) or organisation(s) responsible	AF-NF
=	Public
responsible	
responsible Organisation status	
responsible Organisation status Additional contact	
responsible Organisation status Additional contact Main features	
responsible Organisation status Additional contact Main features Type of database	Public
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biological and histological data that is gathered from morbidly obese patients in standardised conditions and prospectively monitor subjects over a 10-year period. Secondary objective: To gather voluntary subjects to participate in other research projects regarding obesity (impact of bariatric surgery and prognostic factors of its success or failure; research into cardiovascular, metabolic, diagnostic and prognostic factors associated with obesity and its complications, ...)

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Inclusion criteria	$BMI >= 35 \text{ kg/m}^2 \text{ Age: } [18-60] \text{ years old}$

THEIDSION CHICELIA	BMI >= 35 kg/III Age. [10-00] years old
Population type	
Age	Adulthood (19 to 24 years) Adulthood (25 to 44 years) Adulthood (45 to 64 years)
Population covered	Sick population
Gender	Male Woman
Geography area	Regional
French regions covered by the database	Île-de-France
Detail of the geography area	Ile de France

Data collection

Dates

MM/YYYY)

Date of first collection (YYYY or MM/YYYY)	03/2005
Date of last collection (YYYY or	06/2019

Size of the database

Size of the database (number of individuals)	< 500 individuals
Details of the number of individuals	288

Data

Database activity	Current data collection
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Type of data collected	Clinical data Declarative data Paraclinical data Biological data
Clinical data (detail)	Direct physical measures Medical registration
Declarative data (detail)	Paper self-questionnaire Face to face interview
Paraclinical data (detail)	- Knee x-rays - Audio scan Patients who have undergone surgery (bariatric surgery): - OGD endoscopy transit, oesophageal manometry, 24- hour pH monitoring, glucose breath test, liver ultrasound - Ventilatory polygraphy
Biological data (detail)	Blood (systematic dosage): - FBC, haemostasis - Lipid profile - Liver function tests - Fibrinogen, US- CRP - Glucose and insulin level, Renal profile, TSH, DNA (buccal mucosa or blood) Tissue (fat, liver)
Presence of a biobank	Yes
Contents of biobank	Serum Plasma DNA Others
Details of biobank content	Serum bank, plasma bank, DNA bank, histology slides
Health parameters studied	Health event/morbidity Health event/mortality Quality of life/health perception
Procedures	
Data collection method	- Self-administered questionnaire: manual input from a paper questionnaire with double data entry - Interview: manual input from a paper questionnaire with double data entry - Clinical Examinations: handwritten with double data entry - Biological analysis: handwritten with double data entry
Participant monitoring	Yes
Details on monitoring of participants	Duration: 10 years
Links to administrative sources	No
Promotion and access	

Promotion	
Link to the document	http://www.ncbi.nlm.nih.gov/pubmed/23512861
Access	
Terms of data access (charter for data provision, format of data, availability delay)	Data may be used by academic teams or by industrial teams under contract outlining the temporary terms of data usage, according to the decision of the Scientific Committee and AP-HP Department of Clinical Research.
Access to aggregated data	Access on specific project only

Access to individual data

Access on specific project only