ATLANTIS - Long term observational study on the therapeutic management of knee osteoarthritis by general practitioners

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General	
Identification	
Detailed name	Long term observational study on the therapeutic management of knee osteoarthritis by general practitioners
Sign or acronym	ATLANTIS
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	CNIL n°903352 (22/11/2003)
General Aspects	
Medical area	Rheumatology
Others (details)	gonarthrosis, Arthrosis
Keywords	therapeutic management, paracetamol, patient reported outcomes
Scientific investigator(s) (Contact)	
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Unit	Bristol-Myers Squibb
Collaborations	
Funding	
Funding status	Private
Details	Bristol-Myers Squibb
Governance of the database	
Sponsor(s) or organisation(s) responsible	Bristol-Myers Squibb (BMS)
Organisation status	Private
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 care professionals
Database recruitment is carried out as part of an interventional study	No
Additional information regarding sample selection.	Random sampling in clusters
Database objective	
Main objective	Evaluate, in general practice, the change in the first- line therapeutic strategy in patients suffering from knee osteoarthritis, over one year, specially: - Evaluate the percentage of patients treated with each therapeutic modality at the end of the one year observation period. - Describe the modifications implemented in the

therapeutic management and identify the reasons justifying these modifications.

Inclusion criteria

- 1. Males or females 50 years or older suffering from symptomatic osteoarthritis of the knee
- 2. Not treated over the last 6 months for this pathology (non-pharmaceutical treatments and selfmedication by analgesics or NSAIDs at low doses will be allowed).
- 3. For whom the physician will have decided to initiate a first-line treatment in line with the recommendations of EULAR 2000-2003

(paracetamol - systematically taken at a dose of 3 to 4g/d - as pharmacological monotherapy in combination with non-pharmaceutical treatments).

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

International

Detail of the geography area

France and Spain

Data collection

Dates

Date of first collection (YYYY or MM/YYYY)

2003

2006

Date of last collection (YYYY or MM/YYYY)

Size of the database

Size of the database (number of individuals)

[1000-10 000] individuals

Details of the number of

1114 (586 in France, 528 in Spain)

individuals

Data

Database activity	Data collection completed
Type of data collected	Clinical data Declarative data
Clinical data (detail)	Direct physical measures Medical registration
Declarative data (detail)	Face to face interview
Presence of a biobank	No
Health parameters studied	Health care consumption and services Quality of life/health perception
Care consumption (detail)	Hospitalization Medical/paramedical consultation Medicines consumption
Procedures	
Data collection method	paper case report form
Data collection method Participant monitoring	paper case report form Yes
Participant monitoring Details on monitoring of	Yes
Participant monitoring Details on monitoring of participants	Yes M3, M6, M9, M12
Participant monitoring Details on monitoring of participants Links to administrative sources	Yes M3, M6, M9, M12
Participant monitoring Details on monitoring of participants Links to administrative sources Promotion and access	Yes M3, M6, M9, M12
Participant monitoring Details on monitoring of participants Links to administrative sources Promotion and access Promotion	Yes M3, M6, M9, M12

Access on specific project only

Access to individual data