EXFOS - A European prospective observational study to evaluate fracture outcomes, back pain, health-related quality of life, and compliance in patients with osteoporosis during and after treatment with Forsteo®

Head: Médecin pharmacoépidémiologiste, Eli Lilly France

Governance of the database

Last update : 08/01/2012 Version : 2 ID : 79		
General		
Identification		
Detailed name	A European prospective observational study to evaluate fracture outcomes, back pain, health-related quality of life, and compliance in patients with osteoporosis during and after treatment with Forsteo®	
Sign or acronym	EXFOS	
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	CNIL N°910339	
General Aspects		
Medical area	Rheumatology	
Keywords	Osteoporotic patients, teriparatide, duration of treatment, conditions of use	
Scientific investigator(s) (Contact)		
Name of the director	Médecin pharmacoépidémiologiste	
Email	Fr_mail_pharmacoepi@lilly.com	
Unit	Eli Lilly France	
Collaborations		
Funding		
Funding status	Private	
Details	Eli Lilly and Company	

Sponsor(s) or organisation(s) responsible	Eli Lilly
Organisation status	Private
Additional contact	
Main features	
Type of database	
Type of database	Study databases
Study databases (details)	Longitudinal study (except cohorts)
Database recruitment is carried out by an intermediary	A selection of health care professionals
Database recruitment is is made on the basis of:	Medication(s) taken
Database recruitment is carried out as part of an interventional study	No
Additional information regarding sample selection.	France: recruitment by rheumatologists in the course of routine care. Random selection of rheumatologists using professional listing.
Database objective	
Main objective	Primary objective: determine the incidence of clinical vertebral fractures and non-vertebral fragility fractures in patients treated with teriparatide for about 18 months and a post-treatment follow-up period of at least 18 months. Secondary objectives: observance, treatment switching and cessation, clinical evolution, occurrence of back pain, direct costs linked to fractures.
Inclusion criteria	Osteoporotic patients initiating treatment with teriparatide according to the practitioner's opinion in routine care
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	Croatia, Denmark, France, Greece, Italy, Norway, Slovenia and Sweden
Data collection	
Dates	
Date of first collection (YYYY or MM/YYYY)	2006
Date of last collection (YYYY or MM/YYYY)	2012
Size of the database	
Size of the database (number of individuals)	[500-1000[individuals
Details of the number of individuals	1607
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)	Paper self-questionnaire
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	Study data collection form

Participant monitoring	Yes
Details on monitoring of participants	36 months maximum
Links to administrative sources	No
Promotion and access	
Promotion	
Link to the document	http://tinyurl.com/Pubmed-EXFOS
Description	List of publications in Pubmed
Access	
Terms of data access (charter for data provision, format of data, availability delay)	Report and publication
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Access to aggregated data	Access on specific project only