

EDITH - Observational study to evaluate educational modality of type 2 diabetic patients at the time of insulin initiation by GPs

Head :Vignal Franck, Sanofi Aventis

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General	
Identification	
Detailed name	Observational study to evaluate educational modality of type 2 diabetic patients at the time of insulin initiation by GPs
Sign or acronym	EDITH
General Aspects	
Medical area	Endocrinology and metabolism General practice
Others (details)	type-2 diabetes
Keywords	initiation of insulin therapy, insulin
Scientific investigator(s) (Contact)	
Name of the director	Vignal
Surname	Franck
Phone	+33 (0)1 57 63 26 47
Email	franck.vignal@sanofi-aventis.com
Unit	Sanofi Aventis
Collaborations	
Funding	
Funding status	Private
Details	Sanofi-aventis
Governance of the database	
Sponsor(s) or organisation(s) responsible	Sanofi-aventis France

Organisation status	Private
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 selection of health care professionals
Database recruitment 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.	participating general practitioners systematically included the last two consecutive patients that they had put on insulin since less than 6 months and more than 1 month, who met the inclusion criteria and accepted to participate in the study.
Database objective	
Main objective	describe the effects on the fasting glucose at 1 month for main methods of education of type-2 diabetes patients during the initiation of insulin therapy in general practice
Inclusion criteria	man or woman, over the age of 18 years, having a type-2 diabetes responding to the criteria of HAS (French National Authority for Health), treated with oral diabetes medication at the maximum dose (or maximum tolerated dose where applicable), having needed the adding of a basal insulin initiated since at least 1 month and no more than 6 months (+ 1 week) before inclusion, reviewed 1 month after the initiation of insulin, during a routine visit in the framework of following his or her diabetes, accepting to participate in the study after information given by the observing doctor.
Population type	
Age	Adulthood (19 to 24 years)

Adulthood (25 to 44 years)
Adulthood (45 to 64 years)
Elderly (65 to 79 years)

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)	2008
Date of last collection (YYYY or MM/YYYY)	2008
Size of the database	
Size of the database (number of individuals)	[1000-10 000[individuals
Details of the number of individuals	1559
Data	
Database activity	Data collection completed
Type of data collected	Clinical data Biological data
Clinical data (detail)	Direct physical measures
Biological data (detail)	Fasting Blood Glucose
Presence of a biobank	No
Health parameters studied	Health event/morbidity
Procedures	
Data collection method	type-2 diabetes, initiation of insulin therapy, general practice
Participant monitoring	No

Links to administrative sources	No
Promotion and access	
Promotion	
Access	
Terms of data access (charter for data provision, format of data, availability delay)	Poster SFD 2010 methods for accessing the database are currently being defined
Access to aggregated data	Access on specific project only
Access to individual data	Access on specific project only