OPTIMEV - Optimising History Taking for Evaluating The Risk of Venous Thromboembolism

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General		
Identification		
Detailed name	Optimising History Taking for Evaluating The Risk of Venous Thromboembolism	
Sign or acronym	OPTIMEV	
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	N° CCTIRS: 04-459 N° CNIL: 905040 ?05-1118 N° CLINICALTRIALS: NCT00670540	
General Aspects		
Medical area	Cardiology Hematology Pneumology	
Keywords	lower limb deep vein thrombosis, superficial venous thrombosis, post-thrombotic syndrome, pulmonary embolism, Venous thromboembolism, chronic disease, quality of life	
Scientific investigator(s) (Contact)		
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Organization	INSERM - Institut National de la Santé et de la	

Recherche

Name of the director Sevestre

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Organization CHU

Collaborations

Funding

Mixed Funding status

Details Le programme hospitalier de recherche clinique

PHRCSanofi AventisCIC Grenoble et service de

Médecine Vasculaire Grenoble

Governance of the database

Sponsor(s) or organisation(s)

responsible

INSERM - Institut National de la Santé et de la

Recherche Médicale

Organisation status

Sponsor(s) or organisation(s)

responsible

Société Française de Médecine Vasculaire

Organisation status

Public

Sponsor(s) or organisation(s)

responsible

CHU Grenoble

Organisation status

Public

Public

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

A selection of health institutions and services

Database recruitment is carried out as part of an interventional study

No

Additional information regarding sample selection.

Patients referred to a vascular medicine physician for clinically suspected thromboembolism. All patients are enrolled after been seen in one day over 4 recruitment periods (4 seasons)

Database objective

Main objective

To evaluate current risk factors that fit today's practice and long-term prognosis for different anatomical and clinical forms of symptomatic thromboembolism (proximal and distal deep vein thrombosis, superficial venous thrombosis, pulmonary embolism) Follow-up covers recurrent VTE, arterial thrombotic events (coronary, CVA, PAD), onset of cancer, haemorrhagic stroke, postthrombotic syndrome and death.

Inclusion criteria

Adult patients with clinical signs of symptomatic thromboembolism classified after inclusion for VTE+ patients (thromboembolism confirmed through further objective examination) or VTEpatients (thromboembolism ruled out by further examination)

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

National

Detail of the geography area

France, 350 investigating vascular medicine physicians from CHU, CHG, clinics and private

practices

Data collection

Dates

Date of first collection (YYYY or MM/YYYY)

2006

Date of last collection (YYYY or MM/YYYY)

2014

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Size of the database (number of individuals)

[1000-10 000[individuals

Details of the number of individuals

8256

Data

Database activity Current data collection

Type of data collected

Clinical data

Declarative data

Paraclinical data

Clinical data (detail) Direct physical measures

Medical registration

Declarative data (detail) Paper self-questionnaire

Phone interview

Paraclinical data (detail) Doppler ultrasonography and Pulmonary CT

examination report, hospital report

Presence of a biobank No

Health parameters studied Health event/morbidity

Health event/mortality

Quality of life/health perception

Procedures

Data collection method Patient characteristics entered at baseline on an e-

CRF where quality is monitored electronically Followup entered after all target events were validated by a critical events committee (2 physicians evaluating examination and hospital reports) by CRA on an e-

CRF

Participant monitoring Yes

Details on monitoring of

participants

Clinical research associates specialising in vascular disease conducted follow-up telephone interviews with PTS evaluation (Villalta score), quality of life

evaluation as well as events involving venous and arterial thrombosis, cancer, haemorrhagic stroke

and death.

Promotion and access	
Promotion	
Link to the document	http://recherche-clinique.ujf-grenoble.fr/OPTIMEV/
Link to the document	http://tinyurl.com/OPTIMEV
Description	List of publications in HAL
Link to the document	http://www.ncbi.nlm.nih.gov/pubmed/? term=optimev+AND+%28Bosson+JL[Author]+OR +Sevestre+MA[Author]%29+OR+22429908[uid]
Description	List of publications in Pubmed

Access	
Terms of data access (charter for data provision, format of data, availability delay)	Publications
Access to aggregated data	Access on specific project only
Access to individual data	Access on specific project only