SNAC - Study of NorpregnAnes Coagulation

Head:

| Last update : 09/05/2017 Version : 2 ID : 6238 | | |
|--|--|--|
| General | | |
| Identification | | |
| Detailed name | Study of NorpregnAnes Coagulation | |
| Sign or acronym | SNAC | |
| CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation | CCPPRB Bicêtre (21/12/2005), CNIL (19/12/2005) | |
| General Aspects | | |
| Medical area | Biology Endocrinology and metabolism Internal medicine | |
| Pathology (details) | Hormone therapy for menopause | |
| Health determinants | Medicine | |
| Keywords | Oestrogen, progestin, menopause, haemostasis, thrombosis | |
| Scientific investigator(s) (Contact) | | |
| Collaborations | | |
| Funding | | |
| Funding status | Public | |
| Details | Institut National de la Santé et de la Recherche Médicale | |
| Governance of the database | | |
| 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 population file |
| Database recruitment is carried out as part of an interventional study | No |
| Additional information regarding sample selection. | Samples selected on a voluntary basis |
| Database objective | |
| Main objective | Impact of different types of HRT on activated protein C resistance and haemostatic parameters. |
| Inclusion criteria | - Women - between 45 and 70 years of age - menopausal |
| Population type | |
| Age | Adulthood (45 to 64 years) |
| Population covered | General population |
| Gender | Woman |
| Geography area | Regional |
| French regions covered by the database | Île-de-France |
| Detail of the geography area | Centre d'Investigations Préventives et Cliniques de Paris: SS medical examination centre |
| Data collection | |
| Dates | |
| Date of first collection (YYYY or MM/YYYY) | 2006 |
| Date of last collection (YYYY or MM/YYYY) | 2007 |
| Size of the database | |
| Size of the database (number of | < 500 individuals |
| | |

| individuals) | |
|--------------------------------------|--|
| Details of the number of individuals | 300- 100 femmes non traitées/women no treated 100 femmes estrogènes transdermiques + progestérone- 100 femmes estrogènes transdermiques + NOMAC |
| Data | |
| Database activity | Data collection completed |
| Type of data collected | Clinical data Declarative data Paraclinical data Biological data |
| Clinical data (detail) | Direct physical measures |
| Declarative data (detail) | Face to face interview |
| Paraclinical data (detail) | |
| Biological data (detail) | - blood - DNA |
| Presence of a biobank | Yes |
| Contents of biobank | Serum Plasma DNA |
| Details of biobank content | 500 microL plasma aliquots collected with citrate, heparin or EDTA |
| Health parameters studied | Others |
| Other (detail) | |
| Procedures | |
| Data collection method | Standardised questionnaire |
| Participant monitoring | No |
| Links to administrative sources | No |
| Promotion and access | |
| Promotion | |
| Access | |
| Terms of data access (charter | Contact the scientist in charge |

| for data provision, format of data, availability delay) | |
|---|---------------------------------|
| Access to aggregated data | Access on specific project only |
| Access to individual data | Access on specific project only |