- FREGAT

Head: Mariette Christophe

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Identification

Detailed name FREGAT

CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation Accord CPP (10 décembre 2013), ANSM (13 janvier 2014), CCTIRS (12 mars 2014) CNIL (23 décembre 2014)

General Aspects

Medical area Cancer resea	researcl	1
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Health determinants latrogenic

Nutrition

Keywords Tumour of the oesophagus, stomach tumour,

diseases of the oesophagus, digestive system diseases, gastroesophageal cancer, digestive system, clinical and biological records, surgery

Scientific investigator(s)

(Contact)

Name of the director Mariette

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Organization CHRU

Collaborations

Funding

Funding status Public

Details	appel à projet INCa 2012	
Governance of the database		
Sponsor(s) or organisation(s) responsible	CHRU Lille	
Organisation status	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 institutions and services	
Database recruitment is carried out as part of an interventional study	Yes	
Details	Performed at individual level	
Additional information regarding sample selection.	All newly diagnosed patients in participating centres with gastro-oesophageal treatment-naive cancer will be included after acceptance and signed consent, whether they undergo surgery or not, regardless of histological type, tumour stage or therapeutic strategy. Exclusion criteria: - Men or women under the age of 18 People deprived of liberty or under guardianship (including temporary guardianship) People who do not speak French Adults unable to express their consent Patients already included in the FREGAT database Patients who refuse to participate.	
Database objective		
Main objective	Main Objective: To identify, through the creation of a clinico-biological, prospective, multicentric French database, the clinical, biological and tumour factors linked to anti-tumour therapy resistance in patients treated for stage I to IV oesophageal or stomach cancer. Secondary Objectives: - To assess the impact of different current therapeutic strategies for recurrence, survival and quality of life To identify predictive factors for patient treatment	

Inclusion criteria

- Men or women? 18 ans. - Presenting with oesophageal or gastro-oesophageal junction carcinoma recently diagnosed by biopsy, regardless of cancer subtype, tumour stage or proposed treatment. - Treatment-naive cancer. - Have given clear and written consent for blood samples, various questionnaires and the collection of patient information. N.B.: Patients participating in a clinical trial can be included in the FREGAT trial. There is no exclusion period.

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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 Metropolitan France (33 centres, 46 teams)

Data collection

Dates

Date of first collection (YYYY or MM/YYYY)

2014

Size of the database

Size of the database (number of individuals)

Greater than 20 000 individuals

Details of the number of individuals

135 patients au 20/01/2015

Data

Database activity	Current data collection
Type of data collected	Clinical data Declarative data Biological data Administrative data
Clinical data (detail)	Direct physical measures Medical registration
Declarative data (detail)	Paper self-questionnaire
Biological data (detail)	- a collection of tumour samples (pre-treatment biopsies. post-treatment biopsies, surgical specimens) collected in accordance with current quality charters. These samples will be stored in the corresponding investigation centre. Samples will ideally be frozen. However, paraffin conservation will also be acceptable blood sample collection to be put in place in 6 centres with high recruitment potential that have a Biological Resource Centre (CRB) (approximately 3,000 patients). These approved CRBs will ensure sample quality control as per usual.
Administrative data (detail)	The investigating party will issue the patient a FREGAT cohort registration form to be completed by the patient and posted to the Caen Cancéropôle Nord-Ouest Data Processing Centre (CTD/CNO).
Presence of a biobank	Yes
Contents of biobank	Whole blood Serum Plasma Tissues
Details of biobank content	Consult the scientist in charge (Professor Christophe MARIETTE)
Health parameters studied	Health event/morbidity Health event/mortality Health care consumption and services Quality of life/health perception Others
Care consumption (detail)	Hospitalization Medical/paramedical consultation Medicines consumption
Other (detail)	socio-economic questionnaire.

Procedures Data collection method The collection of clinical, biological and epidemiological data as well as the character patient treatment will be stored in the electricase report form (e-CRF) for the study Participant monitoring Yes Details on monitoring of participants Follow-up for 3 years after inclusion. All info in the database is gathered during patient of follow-up, according to usual visit and follow practices in each investigating centre. Annu updates to information is required systemat Updates to information regarding events su death or recurrence will be also required. Links to administrative sources Promotion Link to the document http://www.ncbi.nlm.nih.gov/pubmed/25062 Link to the document http://www.ncbi.nlm.nih.gov/pubmed/25012 Link to the document http://www.ncbi.nlm.nih.gov/pubmed/24919 Link to the document http://www.ncbi.nlm.nih.gov/pubmed/24075 Link to the document http://www.ncbi.nlm.nih.gov/pubmed/23313 Link to the document http://www.ncbi.nlm.nih.gov/pubmed/23064 Link to the document http://www.ncbi.nlm.nih.gov/pubmed/23064		
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Terms of data access (charter for data provision, format of data, availability delay) Publication or presentation of results from the is not permitted without prior agreement from the sponsor (CHRU de Lille) and coordinating researcher (Prof. Christophe MARIETTE).	sion, format of is n y delay) spo	itted without prior agreement from the HRU de Lille) and coordinating
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