## PEPITA - A prospective cohort study assessing the efficacy and safety of Tarceva® in 2nd line in patients with locally advanced or metastatic squamous Non-Small Cell Lung Cancer (NSCLC) - PEPiTA study

Head: Roche Medical Data Center

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General			
Identification			
Detailed name	A prospective cohort study assessing the efficacy and safety of Tarceva® in 2nd line in patients with locally advanced or metastatic squamous Non-Small Cell Lung Cancer (NSCLC) - PEPiTA study		
Sign or acronym	PEPITA		
CNIL registration number, number and date of CPP agreement, AFSSAPS (French Health Products Safety Agency) authorisation	ML28195		
General Aspects			
Medical area	Cancer research		
Study in connection with Covid- 19	No		
Pathology (details)	Non-Small Cell Lung Cancer (NSCLC)		
Health determinants	latrogenic Medicine		
Keywords	ERLOTINIB		
Scientific investigator(s) (Contact)			
Name of the director	Roche Medical Data Center		
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Organization	Roche SAS		

Collaborations	
Participation in projects, networks and consortia	No
Funding	
Funding status	Private
Governance of the database	
Sponsor(s) or organisation(s) responsible	Roche SAS
Organisation status	Private
Presence of scientific or steering committees	Yes
Additional contact	
Name of the contact	https://www.roche.fr/fr/innovation-recherche- medicale/data-sharing-portail-d-information- partage-des-donnees.html
Main features	
Type of database	
Type of database	Study databases
Type of database Study databases (details)	Study databases Cohort study
Study databases (details)  Database recruitment is carried	Cohort study
Study databases (details)  Database recruitment is carried out by an intermediary  Database recruitment is is made	Cohort study  A selection of health institutions and services
Study databases (details)  Database recruitment is carried out by an intermediary  Database recruitment is is made on the basis of:  Database recruitment is carried out as part of an interventional	Cohort study  A selection of health institutions and services  Medication(s) taken

Secondary objectives:

- To describe characteristics of patients treated with Tarceva®:
- To describe Tarceva® treatment modalities in patients with squamous NSCLC after failure of first-line platinum-based chemotherapy;
- To evaluate Overall Survival (OS);
- To describe the evolution of QoL;
- To describe the safety profile of Tarceva®.

### Inclusion criteria

### Inclusion criteria:

- 1. Adult patients (age >= 18 years);
- 2. Histologically and/or cytologically confirmed advanced (stage IIIB) or metastatic (stage IV) NSCLC with predominant squamous component (basaloid carcinomas allowed), after failure of first-line platinum-based chemotherapy;
- 3. For whom the treating physician decided to initiate treatment with Tarceva®:
- 4. Having received oral and written information about the study and having raised no objections to the collection and analysis of his/her personal data.

### Exclusion criteria:

- 1. Mixed non-small cell and small cell lung carcinoma or mixed squamous cell carcinoma with a predominant adenocarcinoma component;
- 2. Concomitant participation in a clinical trial evaluating an antineoplastic treatment.

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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

Pathology

D02 - Carcinoma in situ of middle ear and

respiratory system

Gender

Male Woman

Geography area

National

## Data collection

### **Dates**

Date of first collection (YYYY or MM/YYYY)

2012

Date of last collection (YYYY or MM/YYYY)	2014
Size of the database	
Size of the database (number of individuals)	< 500 individuals
Details of the number of individuals	152
Data	
Database activity	Data collection completed
Type of data collected	Clinical data Biological data
Clinical data (detail)	Medical registration
Details of collected clinical data	Date of visit - Inclusion/eclusion criteria - Demographic data - ECOG - SAP/DAP - Social and occupational data - Risk factors (tobacco use) - Medical history and concomitant diseases - Treatment associated with initiation of Tarceva® - Disease history: primary lung tumor evaluation - Disease history: diagnosis of locally advanced or metastatic disease - EGFR mutation status - First- line treatment of metastatic NSCLC.
Presence of a biobank	No
Health parameters studied	Health event/morbidity Health event/mortality Health care consumption and services Quality of life/health perception
Care consumption (detail)	Medicines consumption
Procedures	
Data collection method	eCRF

# Procedures Data collection method eCRF Classifications used CDISC Quality procedure(s) used GCP/GVP Participant monitoring Yes Monitoring procedures Monitoring by contact with the referring doctor Links to administrative sources No

Promotion and access			
Promotion			
Access			
Dedicated website	https://www.roche.fr/fr/innovation-recherche- medicale/data-sharing-portail-d-information- partage-des-donnees.html		
Presence of document that lists variables and coding procedures	Yes		
Access to aggregated data	Access on specific project only		
Access to individual data	Access on specific project only		