A non-interventional biomarker study in patients with Non-Small Cell Lung Cancer (NSCLC) of adenocarcinoma tumour histology eligible for treatment with Vargatef® according to the approved label (LUME BioNIS)

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## Administrative details

### **EU PAS number**

EUPAS11885

### Study ID

37252

### DARWIN EU® study

No

### **Study description**

To explore whether genetic/genomic markers (alone or combined with clinical covariates) could be used to predict overall survival (OS) in NSCLC patients eligible for treatment with Vargatef® according to the approved label.

### Study status

Finalised

## Research institutions and networks

### Institutions

# Multiple centres: 71 centres are involved in the study

## Contact details

Study institution contact

Boehringer Ingelheim clintriage.rdg@boehringeringelheim.com

Study contact

clintriage.rdg@boehringer-ingelheim.com

**Primary lead investigator** Martin Reck

Primary lead investigator

## Study timelines

Date when funding contract was signed

Actual: 12/06/2015

### Study start date Planned: 09/03/2016 Actual: 09/03/2016

Data analysis start date Planned: 01/11/2019 Actual: 14/03/2016

Date of final study report Planned: 30/06/2020 Actual: 08/07/2020

## Sources of funding

• Pharmaceutical company and other private sector

## More details on funding

Boehringer Ingelheim

## Regulatory

### Was the study required by a regulatory body?

Yes

### Is the study required by a Risk Management Plan (RMP)?

Not applicable

## Methodological aspects

## Study type

## Study type list

### Study topic:

Human medicinal product Disease /health condition

### Study type:

Non-interventional study

Scope of the study:

Other

### If 'other', further details on the scope of the study

Biomarker assessment

### Data collection methods:

Primary data collection

### Main study objective:

Overall Survival in relation to exploratory biomarker assessment, including gene-expressions and genomic alterations.

## Study Design

### Non-interventional study design

Cohort

## Study drug and medical condition

## Name of medicine

### Medical condition to be studied

Non-small cell lung cancer stage IIIB

## Population studied

### Short description of the study population

NSCLC patients eligible for treatment with Vargatef®.

### Age groups

Adults (18 to < 46 years) Adults (46 to < 65 years) Adults (65 to < 75 years) Adults (75 to < 85 years) Adults (85 years and over)

### **Special population of interest**

Other

### Special population of interest, other

Non-small cell lung cancer patients

### Estimated number of subjects

260

## Study design details

### Outcomes

Overall Survival, Disease progression

### Data analysis plan

Biomarkers will be investigated by univariate and multivariate prediction models and regression analysis.For all univariate screening approaches, multiplicity correction of p-values will be performed.Model performances will be quantified and uncertainty will be evaluated by resampling methods.Categorical biomarkers will only be investigated provided there are sufficient patient numbers within the subgroups defined by the categorical biomarkers. For all selected categorical biomarkers the estimate of the HR/odds ratio and its 95% confidence interval (CI) will be presented. For continuous biomarkers the HR/odds ratio for a change per unit will be presented.

## Data management

## Data sources

### Data sources (types)

Other

### Data sources (types), other

Prospective patient-based data collection

## Use of a Common Data Model (CDM)

### CDM mapping

No

## Data quality specifications

### **Check conformance**

Unknown

### **Check completeness**

Unknown

### Check stability

Unknown

### Check logical consistency

Unknown

## Data characterisation

### Data characterisation conducted

No