

## IMREAL - Cohort 2 (dernière mise à jour : 28/06/2019)

[ARCHIMAIDindex.php?action=show&id=706](http://ARCHIMAIDindex.php?action=show&id=706)

### Informations générales

**Titre de l'étude** : A Non-Interventional, Multicenter, Multiple Cohort Study Investigating the Outcomes and Safety of Atezolizumab Under Real-World Conditions in Patients Treated in Routine Clinical Practice

**Traitement** : Métastatique ou localement avancé

**Type d'étude** : Qualité de vie / Observationnelle

**Phase** : NA      **Stade** : Localement avancé à Métastatique      **Ligne(s)** : 2

**Schéma** : This is a non-interventional, multi-country, multi-centre, multiple cohort prospective study, with retrospective collection of prior medical/treatment history data from medical records, designed to assess the real-world outcomes and safety of atezolizumab for indications in the existing label in the real world setting of routine clinical practice.

### Spécialités / Localisations

**Spécialité n°1** : Organes respiratoires et intrathoraciques

**CIM10 - Localisation n°1** : C34 - Tumeur maligne des bronches et du poumon

### Critères

**Critères d'inclusion** : - Patient must have one of the following confirmed diagnoses for which atezolizumab is locally approved in the SmPC:

- (1) As monotherapy for the treatment of adult patients with la/mUC after prior platinum-containing chemotherapy or
- (2) As monotherapy for the treatment of adult patients with la/mNSCLC after prior chemotherapy. Patients with EGFR activating mutations or ALK-positive tumour mutations should also have received targeted therapy before receiving atezolizumab.

- Patient is prescribed atezolizumab therapy for the first time.

- Decision to prescribe atezolizumab must be made and documented prior to inclusion into the study and must follow local clinical practice.

- Initiation of atezolizumab is not more than 28 days before signing informed consent. In countries where enrollment is only allowed after starting treatment, enrollment must be preceded by the administration of the first cycle of atezolizumab.

**Critères de non-inclusion** : - Treatment with atezolizumab for an indication that is not included in the study cohorts. Cisplatin ineligible participants receiving atezolizumab as first line of therapy for la/mUC will be excluded.

- Concomitant anti-cancer therapy at the time of starting atezolizumab on the index date, as per locally approved Summary of Product Characteristics (SmPC).

- Treatment with atezolizumab as part of a clinical trial or for compassionate use as part of an access or compassionate use program.

### Informations promoteur

**Nom du promoteur** : HOFFMANN-LA ROCHE

**Type de promoteur** : Industriel

**Adresse** : - 00000 HORS FRANCE

**Coordonnateur** : - *Mail* : - *Tél* :

## Informations centre investigateur n°1

**Nom du centre** : Clinique Tessier – Groupe AHNAC

**Adresse** : 118 Avenue Désandrouin 59300 VALENCIENNES

**Investigateur** : Docteur Thomas GEY

**TEC / ARC / IDE** : Madame Marielle FERY - *Mail* : fery-m@ch-valenciennes.fr - *Tél* : 03 27 14 07 15

**Ouverture de l'essai** : OUVERT

## Liens utiles

**ClinicalTrials.gov (anglais)** : <https://clinicaltrials.gov/ct2/show/study/NCT03782207>