

Study registered at the country of origin: Specify

Date of registration in national regulatory agency

This ongoing study was submitted before initiation of LBCTR

ASCEND 5: LDK378 Versus Chemotherapy in ALK Rearranged (ALK Positive) Patients Previously Treated With Chemotherapy (Platinum Doublet) and Crizotinib

Protocol number

Type of registration: Justify

Primary sponsor: Country of origin

Novartis Pharmaceuticals

10/11/2014

Acronym

Acronym

CLDK378A2303

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Primary registry identifying number

LBCTR2019121371

MOH registration number

ص/9878

Study registered at the country of origin

Type of registration

Date of registration in national regulatory

10/11/2014

Retrospective

Primary sponsor

Novartis Pharmaceuticals

Date of registration in primary registry

16/05/2023

Public title

ASCEND 5: LDK378 Versus Chemotherapy in ALK Rearranged (ALK Positive) Patients Previously Treated With Chemotherapy (Platinum Doublet) and Crizotinib

Scientific title

A Phase III, Multicenter, Randomized, Open-label Study of Oral LDK378 Versus Standard Chemotherapy in Adult Patients With ALK

-rearranged (ALK-positive) Advanced Non-small Cell Lung Cancer Who Have Been Treated Previously With Chemotherapy (Platinum Doublet) and Crizotinib

Brief summary of the study: English

The primary purpose of the study was to compare the antitumor activity of LDK378 vs. chemotherapy in patients previously treated

with chemotherapy (platinum doublet) and crizotinib.

Brief summary of the study: Arabic

عن طريق الفم مقابل المعالجة الكيميائيّة العاديّة لدى مرضى LDK378 دراسة مرحلة ثالثة متعددة المراكز وجزافيّة ومفتوحة التسمية لدواء بالغين مصابين بسُرطان الرئة غير ذي الخُلايا الصغيرة المتقدّم، كيناز الورم اللّمفي الكشمي المعاد ترتيبه (كيناز الورم اللّمفي الكشمي الإيجابي) وخاضعين سابقًا للمعالجة الكيميائية (البلاتين المزدوج) وللكريز وتينيب

Health conditions/problem studied: Specify

Advanced non-small cell lung cancer (NSCLC)

Interventions: Specify

Drug: Ceritinib

Ceritinib is the investigational treatment and is referred to as the investigational study drug and was provided as 150 mg hard gelatin capsules

for oral use. The dose was 750 mg once daily.





Drug: pemetrexed

Pemetrexed was one of the chemotherapy treatments. Pemetrexed, a reconstituted solution, was intravenously administered over 10 minutes at 500 mg/m2 every 21 days.

Drug: docetaxel

Docetaxel was one of the chemotherapy treatments. Docetaxel, a reconstituted solution, was intravenously administered over 1 hour, at 75 mg/m2 every 21 days.

Key inclusion and exclusion criteria: Inclusion criteria

Inclusion Criteria:

- 1.Patient has a histologically or cytologically confirmed diagnosis of non-small cell lung cancer (NSCLC) that is anaplastic lymphoma kinase (ALK) positive as assessed by the FDA approved Abbott FISH Test.
- 2.Patient has stage IIIB or IV diagnosis and must have received one or two prior regimens (including platinum- doublet) of cytotoxic chemotherapy for the treatment of locally advanced or metastatic NSCLC.
- 3.Patient has at least one measurable lesion as defined by RECIST 1.1. A previously irradiated site lesion may only be counted as a target lesion if there is clear sign of progression since the irradiation
- 4.Patients must have received previous treatment with crizotinib for the treatment of locally advanced or metastatic NSCLC.

Key inclusion and exclusion criteria: Gender

Key inclusion and exclusion criteria: Specify gender

Both

Key inclusion and exclusion criteria: Age minimum

Key inclusion and exclusion criteria: Age maximum

99

N/A

N/A

Key inclusion and exclusion criteria: Exclusion criteria

Exclusion Criteria:

- 1.Patient with known hypersensitivity to any of the excipients of LDK378 (microcrystalline cellulose, mannitol, crospovidone, colloidal silicon dioxide and magnesium stearate)
- 2. Patient with a history of severe hypersensitivity reaction to pemetrexed or docetaxel or any known excipients of these drugs.
- 3.Patient with symptomatic central nervous system (CNS) metastases who is neurologically unstable or has required increasing doses of steroids within the 2 weeks prior to screening to manage CNS symptoms.

Type of study

Interventional

Type of intervention Type of intervention: Specify type

Pharmaceutical

Trial scope Trial scope: Specify scope

Safety

Study design: AllocationStudy design: MaskingRandomized controlled trialOpen (masking not used)

Study design: Control Study phase

Study design: Purpose Study design: Specify purpose

Treatment

Study design: Assignment Study design: Specify assignment

Parallel

IMP has market authorization IMP has market authorization: Specify

Yes, Worldwide

Argentina, Aruba, Australia, Austria, Belgium, Brunei, Canada,
Chile, China, Costa Rica, Croatia, Curacao, Czech Republic,
Denmark, Dominican Republic, El Salvador, Finland, France,

Germany, ...

Name of IMP Year of authorization Month of authorization Month of authorization

LDK378 (Ceritinib)

Type of IMP



Cell therapy

Pharmaceutical class

5-Chloro-2-N-{5-methyl-4-(piperidin-4-yl)-2-[(propan-2-yl)oxy]phenyl}-4-N-[2-(propane-2-sulfonyl) phenyl]pyrimidine-2,4-diamine

Therapeutic indication

This study will be conducted in adult male or female patients, with ALK-rearranged (as determined by the Abbott FISH test), advanced (Stage IIIB or IV) NSCLC, who have received previous treatment with cytotoxic chemotherapy (one or two prior regimens, including one platinum doublet) and crizotinib, and have demonstrated disease progression at study enrollment. No particular sequence of prior crizotinib and chemotherapy is required for enrollment, and either can comprise the last treatment received by the patient.

Therapeutic benefit

Progression Free Survival (PFS) and Overall Survival (OS)

Study model Study model: Explain model

N/A N/A

Study model: Specify model

N/A

Time perspective Time perspective: Explain time perspective

NA

N/A

Time perspective: Specify perspective

Target follow-up duration Target follow-up duration: Unit

Number of groups/cohorts

Biospecimen retention Biospecimen description

None retained

Target sample size Actual enrollment target size

Date of first enrollment: Type Date of first enrollment: Date

28/01/2015 Actual

Date of study closure: Date Date of study closure: Type

Actual 04/04/2023

Recruitment status: Specify Recruitment status

Complete





Date of completion

30/10/2015

IPD sharing statement plan

No

IPD sharing statement description

Novartis is committed to sharing with qualified external researchers, access to patient-level data and supporting clinical documents from eligible studies. These requests are reviewed and approved by an independent review panel on the basis of scientific merit. All data provided is anonymized to respect the privacy of patients who have participated in the trial in line with applicable laws and regulations.

This trial data availability is according to the criteria and process described on www.clinicalstudydatarequest.com

Additional data URL

Admin comments

Trial status

Approved

Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
clinicaltrials.gov	NCT01828112	

Sources of Monetary or Material Support

Name

Novartis Pharmaceuticals

Secondary Sponsors

Name

NA

Contac	Contact for Public/Scientific Queries						
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation	
Public	Marwan Ghosn	Beirut	Lebanon	03-226842	marwanghosnmd @yahoo.com	Hotel Dieu De France	
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Centers/Hospitals Involved in the Study			
Center/Hospital name	Name of principles investigator	Principles investigator speciality	Ethical approval
Hotel Dieu De France	Marwan Ghosn	Hematology oncology	Approved

Ethics Review					
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone	
Hotel Dieu de France	22/10/2014	Sami Richa	cue@usj.edu.lb	961421229	

Countries of Recruitment
Name
Lebanon
Belgium
France
Canada
Germany
Italy
Japan
Netherlands
Turkey
United Kingdom
United States of America

	Health Conditions or Problems Studied				
Condition		Code	Keyword		
	Advanced non-small cell lung cancer (NSCLC)	Bronchus or lung, unspecified (C34.9)	Advanced non-small cell lung cancer (NSCLC)		



Interventions				
Intervention	Description	Keyword		
ICF, physical assessment, ECG, radiology, PK sampling	ICF, physical assessment, ECG, radiology, PK sampling	ICF, physical assessment, ECG, radiology, PK sampling		

Primary Outcomes				
Name	Time Points	Measure		
Progression Free Survival (PFS)	24 months	24 months		

Key Secondary Outcomes				
Name	Time Points	Measure		
Overall Survival (OS)	18 months	18 months		
Overall Response Rate (ORR) [18 months	18 months		
Patient Reported Outcomes (PRO)	every 6 weeks	every 6 weeks		



Trial Results	
Summary results	
Study results globally	
Date of posting of results summaries	Date of first journal publication of results
Results URL link	
Baseline characteristics	
Participant flow	
Adverse events	
Outcome measures	
URL to protocol files	