PATIENT
EXAMPLE-LUNG-01
REPORT DATE
17-Sep-2025

Gender: Female | Birth year: 1975 | WHO: 1

Tumor: Lung adenocarcinoma | Lesions: Liver, Lung | Stage: IV

### **Clinical summary**

Relevant systemic treatment history 6/2023-1/2025 Osimertinib

#### Recent molecular results

#### Hartwig WGS (22-Feb-2025)

Biopsy location Lung (purity 50%)

Molecular tissue of origin prediction Lung: Non-small cell: LUAD (98%)

Tumor mutational load / burden TML 160 / TMB 14 mut/Mb

Microsatellite (in)stability Stable

HR status Proficient (0)

Driver mutations EGFR C797S, EGFR L858R, KRAS G12C, KRAS G12D

Amplified genes None

Deleted genes TP53

Homozygously disrupted genes None

Gene fusions MET(exon13)::MET(exon15) fusion

Virus None

**Trial-relevant IHC results** 

PD-L1 Score > 50%

### Trials in NL that are open and potentially eligible (5 cohorts from 5 trials)

Trial	Cohort	Molecular	Sites	Warnings
METC 04 TEDR1	Lung cancer C797S cohort	EGFR C797S	NKI-AvL	None
METC 02 KAYRAS	Dose expansion - monotherapy - NSCLC	KRAS G12D, PD-L1 >= 50.0	Erasmus MC	Variant(s) G12D in KRAS but subclonal likelihood of > 50%
METC 01 IEMOEN	Dose escalation - monotherapy (no slots)	None		Has not exhausted SOC (at least platinum doublet remaining)
EGFR-C797S- TRIAL	EGFR C797S	EGFR C797S	Elisabeth- TweeSteden Ziekenhuis	
EGFR-L858R- TRIAL	EGFR L858R	EGFR L858R	Elisabeth- TweeSteden Ziekenhuis	

Trials matched solely on molecular event and tumor type (no clinical data used) are shown in italicized, smaller font.

### International trials that are open and potentially eligible (1 cohort from 1 trial)

Trial	Cohort	Molecular	Sites	
KRAS-G12C-TRIAL-DE	KRAS G12C	KRAS G12C	Germany: Stuttgart	

International trials are matched solely on molecular event and tumor type (clinical data excluded).

<sup>1</sup> trial filtered due to trials recruiting nationally for the same molecular target. See Other Trial Matching Results for filtered matches.

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### **Molecular Details**

### Hartwig WGS (EXAMPLE-LUNG-01-T, 22-Feb-2025)

#### General

Purity	Ploidy	TML Status	TMB Status	MS Stability	HR Status	DPYD	UGT1A1
- unity	Fioldy	TIVIL Status	TWD Status	WIS Stability	Titi Status	DF 1D	OGITAL
50%	2.3	High (160)	High (14)	Stable	Proficient (0)	*1_HOM (Normal function)	*1_HOM (Normal function)
Predicted tumo	or origin						
				1. Lung: Non-sma	all cell: LUAD		
Combined prediction score			98%				
This score is ca	alculated by combini	ng information on:					
(1) SNV ty	pes			60%			
(2) SNV ge	enomic localisation	distribution		70%			
(3) Driver (	genes and passeng	er characteristics		80%			

Other cohorts have a combined prediction of 2% or lower

### **Key drivers**

Туре	Driver	Trials (Locations)	Trials in Hartwig	Best evidence in External	Resistance in External
Mutation (gain of function)	EGFR L858R (2/4 copies)		NCT00000006	Approved	
Mutation (gain of function)	EGFR C797S (1/4 copies)	TEDR1 (NKI-AvL)	NCT00000008	Pre-clinical	
Mutation (gain of function)	KRAS G12D (0.3/2 copies)*	KAYRAS (Erasmus MC)			
Mutation (gain of function)	KRAS G12C (0.3/2 copies)*		NCT00000009		
Deletion	TP53 del, 0 copies				
Known fusion	MET(exon13)::MET(exon15) fusion				

<sup>\*</sup> Variant has > 50% likelihood of being sub-clonal

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Other drivers or relevant events

Type Driver Trials (Locations) Trials in Hartwig Best evidence in Resistance in External External

None

**IHC** results

PD-L1 Score > 50%

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### **Clinical Details**

### **Clinical summary**

Relevant systemic treatment history 6/2023-1/2025 Osimertinib

Relevant other oncological history None

Previous primary tumor None

Relevant non-oncological history 2023 Rheumatoid arthritis

Patient current details (20-Feb-2025)

Unresolved toxicities grade => 2

LVEF

50%

Known allergies

None

Recent surgeries 01-Aug-2024 Cholecystectomy

Tumor details (20-Feb-2025)

Measurable disease Yes

Known lesions Liver, Lung
Unknown lesions None

No lesions present CNS, Brain, Bone, Lymph node

### **Active medication details**

Medication	Administration route	Start date	Stop date	Dosage	Frequency
St. John's Wort	Oral	01-Feb-2023		300 MILLIGRAMS	1 / 2 DAYS

## **Blood transfusions**

Product	Date
ERTHROCYTES_FILTERED	20-Sep-2024

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## **Trial Matching Details**

Filtered trials potentially eligible based on molecular results which are potentially recruiting (1)

Trial	Cohort	Molecular	Sites
<u>EGFR-BE</u>	EGFR L858R	EGFR L858R	Belgium: Brussels

### Trials and cohorts that are potentially eligible, but are closed (1)

Trial	Cohort	Molecular	Sites	Warnings
METC 01 IEMOEN	Dose expansion - monotherapy	None		Has not exhausted SOC (at least platinum doublet remaining)

### Trials and cohorts that are considered ineligible (4)

Trial	Cohort	Molecular	Ineligibility reasons
METC 03 NO-SEE797ES	Dose escalation - monotherapy	EGFR C797S	C797S in EGFR in canonical transcript
METC 02 KAYRAS	Dose expansion - monotherapy - Colorectum	KRAS G12D, PD-L1 >= 50.0	No colorectal cancer
METC 05 PICKME3CA	Applies to all cohorts below	None	No PIK3CA activating mutation(s)
	Dose expansion - monotherapy - NSCLC (closed)		
	Dose expansion - monotherapy - Other cancer types (closed)		Tumor belongs to DOID term(s) lung non-small cell carcinoma

### Trials and cohorts that are not evaluable or ignored (0)

Trial	Cohort	Molecular	Sites	Configuration

None