PATIENT
EXAMPLE-LUNG-01
REPORT DATE
07-Nov-2024

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

Tumor: Lung - Adenocarcinoma | Lesions: Liver, Lung | Stage: IV

## **Summary**

### **Clinical summary**

Gender Female Birth year 1975

WHO 1 Tumor Lung - Adenocarcinoma

Lesions Liver, Lung Stage IV

Measurable disease Yes DPYD \*1\_HOM (Normal function)

(RECIST)

UGT1A1 \*1\_HOM (Normal function)

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

Relevant other oncological history

Previous primary tumor

None

Relevant non-oncological history 2023 Rheumatoid arthritis

Recent molecular results KRAS G12C (0.3/2 copies)\*, KRAS G12D (0.3/2 copies)\*, NRAS: No reportable

events, BRAF: No reportable events, HER2: No reportable events, MSS

#### **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 High (160) / TMB High (14)

Microsatellite (in)stability Stable

HR status Proficient (0)

High driver mutations EGFR L858R, EGFR C797S, KRAS G12D, KRAS G12C

Amplified genes

Deleted genes

TP53

Homozygously disrupted genes

Gene fusions

Virus detection

Trial-relevant events, considered medium/low driver:

None

IHC results PD-L1: Score > 50%

## Standard of care options considered potentially eligible

There are no standard of care treatment options for this patient

### Approved treatments considered eligible

**Treatment** 

Not yet determined

### Trials in NL that are open and potentially eligible (4 cohorts from 4 trials)

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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	Erasmus MC	Variant(s) G12D in KRAS but subclonal likelihood of > 50%
EGFR-C797S-TRIAL	EGFR C797S	EGFR C797S	Elisabeth-TweeSteden Ziekenhuis	
EGFR-L858R-TRIAL	EGFR L858R	EGFR L858R	Elisabeth-TweeSteden Ziekenhuis	

# International trials that are open and potentially eligible (2 cohorts from 2 trials)

Trial	Cohort	Molecular	Sites
EGFR-BE	EGFR L858R	EGFR L858R	Belgium (Brussels)
KRAS-G12C-TRIAL-DE	KRAS G12C	KRAS G12C	Germany (Stuttgart)

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

## Trials and cohorts that are considered ineligible (2)

Trial	Cohort	Molecular Sites	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 Erasmus MC G12D	No colorectal cancer

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# Resistance evidence

		evi		

There are no standard of care treatment options for this patient

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## **Molecular Details**

**IHC results** PD-L1: Score > 50%

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

Other cohorts have a combined prediction of 2% or lower

General							
Purity	Ploidy	TML Status	TMB Status	MS Stability	HR Status	DPYD	UGT1A1
50%	2.3	High (160)	High (14)	Stable	Proficient (0)	*1_HOM (Normal function)	*1_HOM (Normal function)
Predicted	l tumor origir	1					
						1. Lung: Non-small cell	: LUAD
Combine	ed prediction	score				98%	
This scor	e is calculated	by combining inform	nation on:				
(1) 5	SNV types					60%	
(2) \$	SNV genomic	localisation distribution	on			70%	
(3)	Oriver genes a	nd passenger charac	eteristics			80%	

#### **Drivers**

Туре	Driver	Driver likelihood	Trials (Locations)	Trials in Hartwig	Best evidence in External	Resistance in External
Mutation (Hotspot)	EGFR C797S (1/4 copies)	High	TEDR1 (NKI-AvL)	NCT00000008	Pre-clinical	
Mutation (Hotspot)	EGFR L858R (2/4 copies)	High		NCT00000006, NCT00000007	Approved	
Mutation (Hotspot)	KRAS G12C (0.3/2 copies)*	High		NCT00000009		
Mutation (Hotspot)	KRAS G12D (0.3/2 copies)*	High	KAYRAS (Erasmus MC)			
Loss	TP53 del, 0 copies	High				
Known fusion	MET::MET, exon 13 - exon 15	High				
* Variant has > 50% like	lihood of being sub-clonal					

All results and data described in this report are for Research Use Only and have NOT been generated using a clinically validated and controlled procedure nor is it a validated medical device. The results should NOT be used for diagnostic or treatment purposes. No rights can be derived from the content of this report. 4/10 Gene and variant annotations and related content are powered by Genomenon Cancer Knowledgebase (CKB).

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# **Molecular History**

Molecular history				
Event	Description	Driver likelihood	2025-02-22	
			Hartwig WGS	
EGFR L858R	Mutation (Hotspot)	High	VAF 0.5%	
(Tier I)	Gain of function			
EGFR C797S	Mutation (Hotspot)	High	VAF 0.25%	
(Tier II)	Gain of function			
KRAS G12C	Mutation (Hotspot)	High	VAF 0.15%	
(Tier III)	Gain of function			
KRAS G12D	Mutation (Hotspot)	High	VAF 0.15%	
(Tier III)	Gain of function			
MET::MET	Known fusion	High	Detected	
(Tier III)	Gain of function			
TP53 del	Loss	High	Detected	
(Tier III)	Unknown protein effect			
ТМВ			14.0	
MSI			Stable	

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# **SOC literature efficacy evidence**

## Standard of care options considered potentially eligible

The following standard of care treatment(s) could be an option for this patient. For further details per study see 'SOC literature details' section in extended report.

There are no standard of care treatment options for this patient

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## **Clinical Details**

ent history	6/2023-1/2025	Osimertinib		
I history	None			
	None			
history	2023	Rheumatoid arthri	tis	
(20-Feb-2025)				
e => 2	None			
	50%			
ons	None			
	None			
	01-Aug-2024 Cl	nolecystectomy		
-2025)				
	Yes			
	No known CNS	lesions		
	No known brai	n lesions		
ails				
dministration route	Start date	Stop date	Dosage	Frequency
)ral	01-Feb-2023		300 MILLIGRAMS	1 / 2 DAYS
			Date	
ED			20-Sep-2024	
	ent history I history (20-Feb-2025) e => 2 ons -2025)  ails administration route Oral	None	None None None None None (20-Feb-2025)  e => 2  None None None None None O1-Aug-2024 Cholecystectomy  -2025)  Yes No known CNS lesions No known brain lesions  ails ails administration route Start date Oral O1-Feb-2023	None None history  2023 Rheumatoid arthritis  (20-Feb-2025)  e => 2 None 50% ons None None 01-Aug-2024 Cholecystectomy  -2025)  Yes No known CNS lesions No known brain lesions  ails  dministration route Start date O1-Feb-2023 Sou MILLIGRAMS

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## **SOC literature details**

There are no standard of care treatment options for this patient

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# **Molecular Evidence**

On label clinical	evidence					
Event	CKB Event	Level A	Level B	Level C	Level D	
EGFR C797S	EGFR C797S				AFATINIB	
					Lung non-small o	cell carcinoma (2015)
EGFR L858R	EGFR L858R	OSIMERTINIB				
		Lung non-small cell carcinoma (2016)				
		AFATINIB				
		Lung non-small cell carcinoma (2013)				
Off label clinical	evidence					
Event	CKB Event	Level A	Level B	Leve	el C	Level D
None						
Efficacy evidence	e description					
EGFR L858R						
OSIMERTINIB:		Level A (2016)	Lung non-small cell ca	rcinoma	Osimertinib	is effective in patients with R mutations
AFATINIB:		Level A (2013)	Lung non-small cell ca	rcinoma	Afatinib is ef L858R muta	fective in patients with EGFR tions
EGFR C797S						
AFATINIB:		Level D (2015)	Lung non-small cell ca	rcinoma		port, afatinib was effective R L858R/C797S positive lung

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# **Trial Matching Overview**

None

	Cohort	Molecular	Sites	Warnings	
EGFR-C797S-TRIAL	EGFR C797S	EGFR C797S	Elisabeth-TweeStede Ziekenhuis	en	
EGFR-L858R-TRIAL	EGFR L858R	EGFR L858R	Elisabeth-TweeStede Ziekenhuis	en	
Trials matched solely on	molecular event and tumor type (no clir	nical data used) are	e shown in italicized, sr	maller font.	
International trials	that are open and potentially	y eligible (2 co	horts from 2 tria	ıls)	
Trial	Cohort			Molecular	Sites
EGFR-BE	EGFR L858R			EGFR L858R	Belgium (Brussels)
KRAS-G12C-TRIAL-E	KRAS G12C			KRAS G12C	Germany (Stuttgart)
		mor typo (olinical o	lata excluded)		
International trials are ma	atched solely on molecular event and tu	illioi type (ciillicai c	ala onoladoa).		