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 1/2023-9/2024 Osimertinib

Relevant other oncological history

Previous primary tumor

None

Relevant non-oncological history 2022 Rheumatoid arthritis

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

events, HER2: No reportable events, MSS

Recent molecular results

Hartwig WGS (01-Sep-2024)

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

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

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

| Trial | Cohort | Molecular | Warnings |
|-------------------|--------------------------------|------------|--|
| METC 04 TEDR1 | Lung cancer C797S cohort | EGFR C797S | None |
| METC 02 KAYRAS | Dose expansion - monotherapy - | KRAS G12D | Variant(s) KRAS G12D in KRAS but subclonal likelihood of > 50% |

Example trials that are open and potentially eligible but currently have no slots available (1 cohort from 1 trial)

| Trial | Cohort | Molecular | Warnings |
|---------|-------------------------------|-----------|---|
| METC 01 | Dose escalation - monotherapy | | History of Rheumatoid arthritis, SOC not exhausted: at least platinum |
| IEMOEN | | | doublet remaining |

External trials potentially eligible based on molecular results which are potentially recruiting locally in Netherlands (2)

| Trial title | Events | Source Events | Cancer Types | Hospitals |
|------------------|---------------------------|---------------------------|-------------------------------|--------------------------|
| EGFR-BE | EGFR C797S, EGFR L858R | EGFR C797S, EGFR L858R | Lung non-small cell carcinoma | Elisabeth- TweeSteden |
| | | | | Ziekenhuis |
| EGFR-L858R-TRIAL | EGFR L858R | EGFR L858R | Lung non-small cell | Elisabeth- |
| | | | carcinoma | TweeSteden |
| | | | | Ziekenhuis |

Example trials and cohorts that are open but considered ineligible (4)

| Trial | Cohort | Molecular | Ineligibility reasons |
|------------------------|---|-----------|---|
| METC 02 KAYRAS | Dose expansion - monotherapy - Colorectum | KRAS G12D | No colorectal cancer |
| METC 03 NO-SEE797ES | Dose escalation - monotherapy | | C797S detected in EGFR |
| METC 05 PICKME3CA | Applies to all cohorts below Dose expansion - monotherapy - NSCLC | None | No PIK3CA activating mutation(s) |
| | Dose expansion - monotherapy - Other cancer types | | Has tumor belonging to DOID term(s) lung non-small cell carcinoma |

Open cohorts with no slots available are shown in grey.

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Resistance evidence

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, 01-Sep-2024)

General

| Purity | TML Status | TMB Status | MS Stability | HR Status | DPYD | UGT1A1 |
|--------|------------|------------|--------------|----------------|--------------------------|--------------------------|
| 50% | High (160) | High (14) | Stable | Proficient (0) | *1_HOM (Normal function) | *1_HOM (Normal function) |

Predicted tumor origin

1. Lung: Non-small cell: LUAD

| | _ |
|---|-----|
| Combined prediction score | 98% |
| This score is calculated by combining information on: | |
| (1) SNV types | 60% |
| (2) SNV genomic localisation distribution | 70% |
| (3) Driver genes and passenger characteristics | 80% |

Other cohorts have a combined prediction of 2% or lower

Drivers

| Туре | Driver | Driver likelihood | Trials in Example | Trials in Hartwig | Best evidence in External | Resistance in External |
|--------------------|----------------------------|-------------------|-------------------|-----------------------------|------------------------------|------------------------|
| Mutation (Hotspot) | EGFR C797S (1/4 copies) | High | TEDR1 | NCT0000007 | Pre-clinical | |
| Mutation (Hotspot) | EGFR L858R (2/4 copies) | High | | NCT00000007, NCT00000006 | Approved | |
| Mutation (Hotspot) | KRAS G12D (0.3/2 copies)* | High | KAYRAS | | | |
| Loss | TP53 del, 0 copies | High | | | | |
| Known fusion | MET_MET, exon 14 - exon 14 | High | | | | |

^{*} Variant has > 50% likelihood of being sub-clonal

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Molecular History

Molecular history

| Event | Description | Driver likelihood | 2024-09-01 Hartwig WGS |
|-------------------------|--|-------------------|---------------------------|
| EGFR L858R (Tier I) | Missense Gain of function Hotspot | High | VAF 0.5% |
| EGFR C797S (Tier II) | Missense Gain of function Hotspot | High | VAF 0.25% |
| KRAS G12D (Tier III) | Missense Gain of function Hotspot | High | VAF 0.15% |
| MET_MET (Tier III) | Fusion Known fusion Gain of function | High | Detected |
| TP53 del (Tier III) | Deletion Unknown protein effect | High | Detected |
| 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

Clinical summary

Relevant systemic treatment history 1/2023-9/2024 Osimertinib

Relevant other oncological history None

Previous primary tumor None

2022 Rheumatoid arthritis Relevant non-oncological history

Patient current details (01-Oct-2024)

Unresolved toxicities grade => 2 None 50% **LVEF** Cancer-related complications None Known allergies None

Recent surgeries 01-Aug-2024 Cholecystectomy

Tumor details (01-Oct-2024)

Measurable disease Yes

CNS lesion status No known CNS lesions Brain lesion status No known brain lesions

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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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 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 Level C Level D

Efficacy evidence description

EGFR L858R

OSIMERTINIB: Level A (2016) Lung non-small cell carcinoma Osimertinib is effective in patients with EGFR L858R mutations

AFATINIB: Level A (2013) Lung non-small cell carcinoma Afatinib is effective in patients with EGFR L858R mutations

EGFR C797S

AFATINIB: Level D (2015) Lung non-small cell carcinoma In a case-report, afatinib was effective against EGFR L858R/C797S positive lung cancer.

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Trial Matching Summary

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| EGFR-L858R-TRIAL | EGFR L858R | EGFR L858R | Lung non-small cell carcinoma | Elisabeth- TweeSteden Ziekenhuis |