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

Gender: Female | Birth year: 1946 | WHO: 0

Tumor: Colorectum (cecum) carcinoma | Lesions: Lung, Peritoneal | Stage: IV

### **Clinical summary**

Relevant systemic treatment history None

Relevant other oncological history 11/2021 Hemicolectomy right (Cecum)

Previous primary tumor Skin squamous cell carcinoma (diagnosed 6/2016, last treatment 8/2016, considered

non-active)

Relevant non-oncological history 1/2019 Cerebrovascular accident

### Standard-of-care options considered potentially eligible

Treatment	Literature efficacy evidence	Real-world efficacy evidence Warnings	Warnings	
FOLFIRI	PHASE-3-CRC			
	<b>PFS</b> : 10.0 months (95% CI: 10.0-12.0)	<b>PFS:</b> 13.3 months, IQR: 12.6		
	<b>OS:</b> 25.0 months (95% CI: 25.0-30.0)	<b>OS:</b> 22.2 months, IQR: 24.8		

### Trials in NL that are open and potentially eligible (1 cohort from 1 trial)

Trial	Cohort	Molecular	Sites	Warnings
KRAS-G12D-	KRAS G12D	KRAS G12D	UMC Utrecht	
<u>TRIAL</u>				

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

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## **Molecular Details**

## **NGS & MSI Panel (15-Jan-2023)**

Biopsy location Lung

Tumor mutational burden TMB 8 mut/Mb

Microsatellite (in)stability Stable

Driver mutations KRAS G12D

**IHC** results

Ki67 Positive, score 90%

PD-L1 Score < 50%

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## **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.

Treatment	Literature efficacy evider	nce
FOLFIRI	PHASE-3-CRC	
	Patient characteristics:	
	WHO/ECOG	0: 100, 1: 80, 2: 20, 3: 0, 4: 0
	Primary tumor location	Left: 145, Both or unknown: 10, Right: 45
	Mutations	KRAS exon 2 wild-type 200/200
	Metastatic sites	Liver only: 58 (32.0%), Lung only: 10 (6.0%)
	Previous systemic therap	by 35/200
	Prior therapies	Adjuvant chemotherapy
	Median PFS:	10.0 months (95% CI: 10.0-12.0)
	Median OS:	25.0 months (95% CI: 25.0-30.0)

#### Treatment decisions (percentage of population assigned to systemic treatment) in NCR real-world data set

	All (n=9207)	Age 73-83y (n=2727)	WHO 1 (n=2828)	RAS positive (n=2760)	Liver only lesions (n=2715)
FOLFIRI	38.5%	23.8%	37.9%	44.6%	39.5%
Median overall survival (OS)	in months in NCR real-world data	set			
	All (n=9207)	Age 73-83y (n=2727)	WHO 1 (n=2828)	RAS positive (n=2760)	Liver only lesions (n=2715)
FOLFIRI	<b>16.1</b> , IQR: 18.2	<b>15.4</b> , IQR: 18.2	<b>14.8</b> , IQR: 16.3	<b>15.8</b> , IQR: 14.2	<b>16.5</b> , IQR: 17.4
	(n=3543)	(n=649)	(n=1071)	(n=1230)	(n=1073)

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#### Median progression-free survival (PFS) in months in NCR real-world data set

	All (n=5018)	Age 73-83y (n=1330)	WHO 1 (n=1623)	RAS positive (n=1822)	Liver only lesions (n=1534)
FOLFIRI	<b>8.2</b> , IQR: 5.5	<b>8</b> , IQR: 6.1	<b>7.9</b> , IQR: 5	<b>8</b> , IQR: 4.7	<b>8.3</b> , IQR: 5.3
	(n=2106)	(n=340)	(n=661)	(n=836)	(n=652)

#### **Explanation:**

These tables only show treatments that are considered standard of care (SOC) in colorectal cancer in the Netherlands.

The 'All' column shows results in NCR patients who were previously untreated, diagnosed with colorectal cancer with distant metastases and treated systemically without surgery, for whom the treatment could be categorized in SOC treatments.

The 'Age', 'WHO', 'RAS' and 'Lesions' columns show results based on patients from the 'All' population, filtered for equal WHO, similar age, equal RAS status or equal lesion localization, respectively.

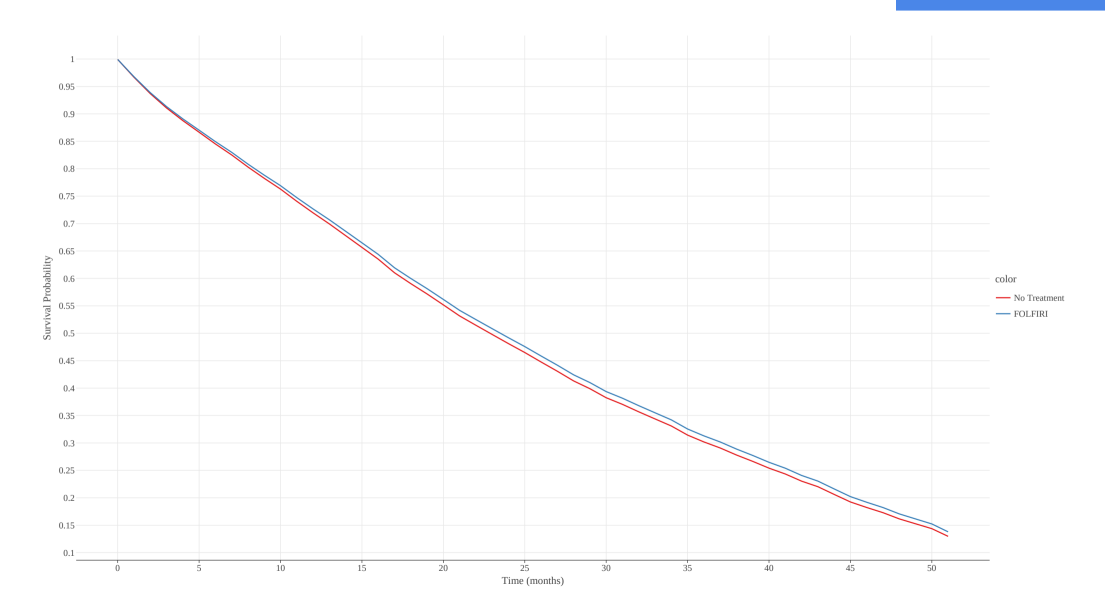
'PFS' is calculated as the duration from the date on which the first compound of the treatment was administered, until first progression.

'OS' is calculated as the duration from the date on which the first compound of the treatment was administered, until death from any cause.

When patient number is too low (n  $\leq$  20) to predict PFS or OS, "NA" is shown.

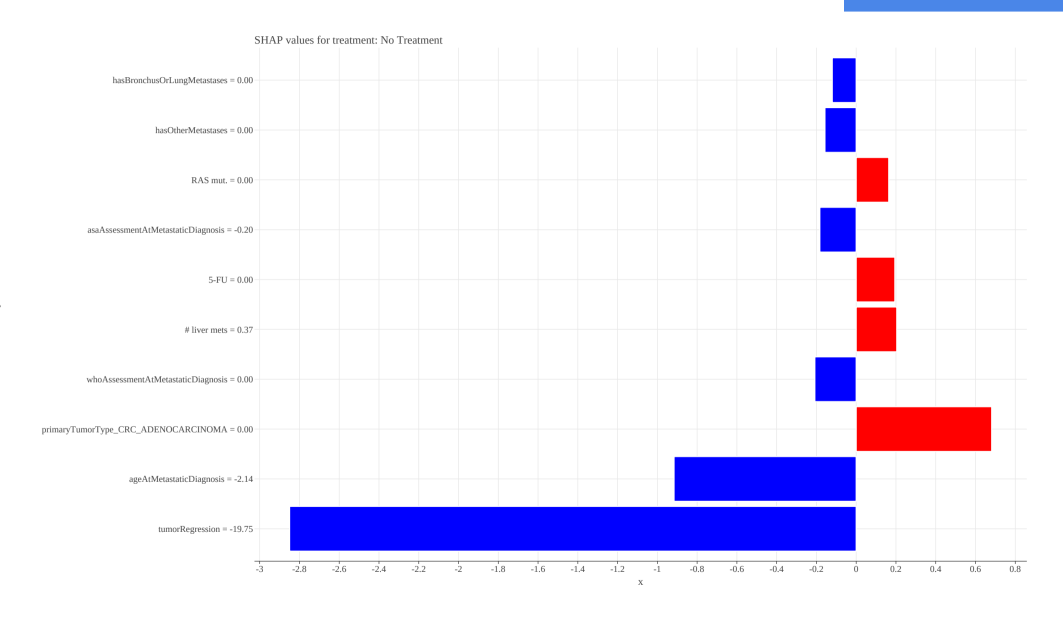
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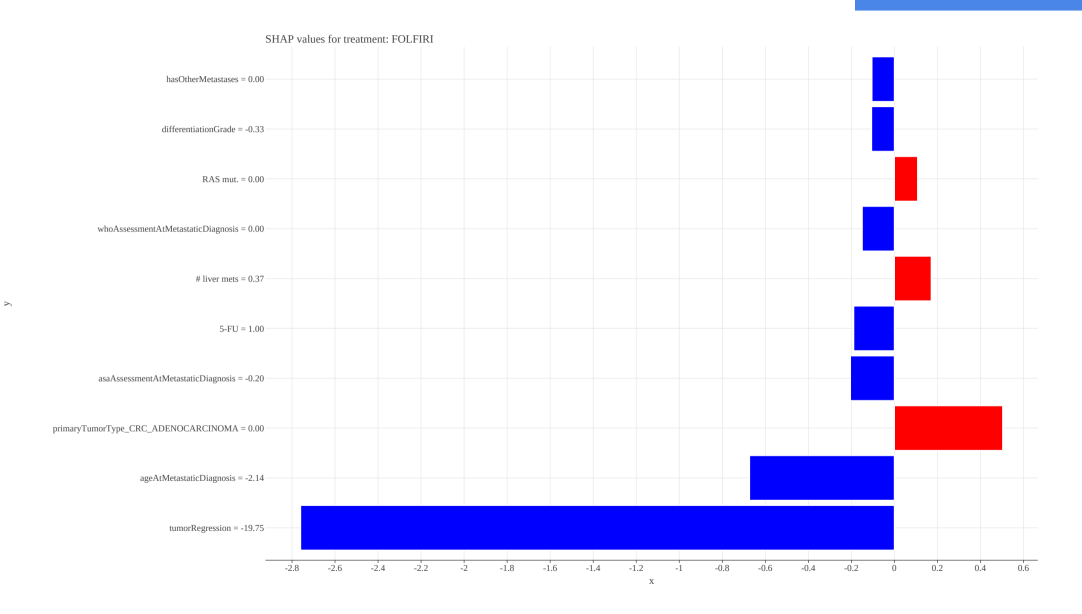


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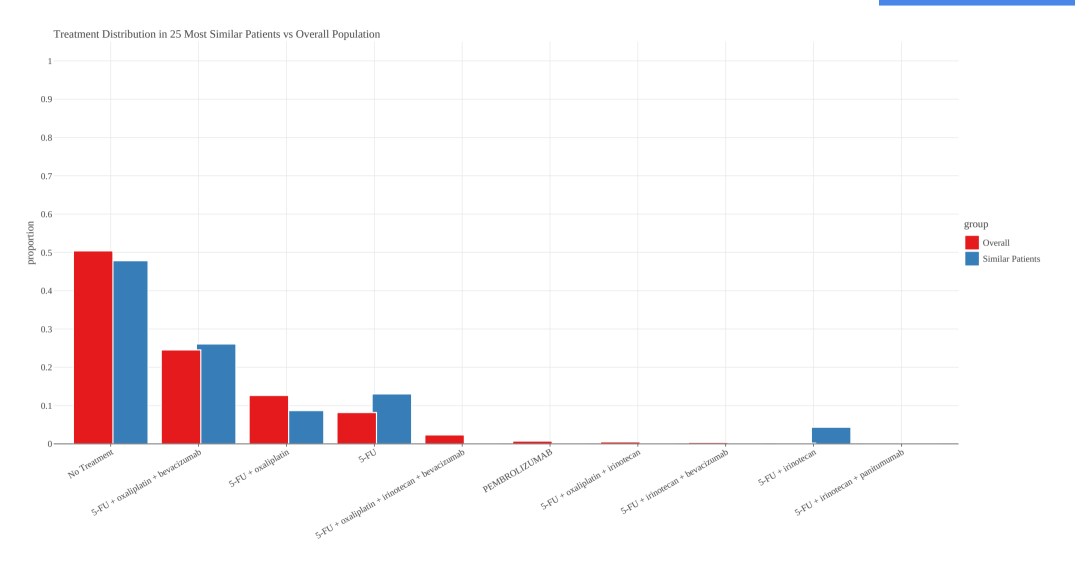


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treatment

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## Resistance evidence

Treatment	Mutation	Evidence source	Evidence level	Found in molecular analysis
FOLFIRI	GENE S11C	[1]	D	Yes

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### **Clinical Details**

## **Clinical summary**

Relevant systemic treatment history None

Relevant other oncological history 11/2021 Hemicolectomy right (Cecum)

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Relevant non-oncological history 1/2019 Cerebrovascular accident

Patient current details (05-Mar-2023)

Unresolved toxicities grade => 2

Cancer-related complications

Known allergies

Morphine

Recent surgeries 12-Nov-2021 Hemicolectomy right

Tumor details (05-Mar-2023)

Measurable disease Yes

Known lesions Lung, Peritoneal Unknown lesions Lymph node

No lesions present CNS, Brain, Liver, Bone

**Active medication details** 

Medication Administration route Start date Stop date Dosage Frequency

None

**Blood transfusions** 

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ERTHROCYTES\_FILTERED 10-Jan-2023

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## **Trial Matching Details**

Trials and cohorts that are potentially eligible, but are closed (2)

Trial	Cohort	Molecular	Sites	Warnings
METC 01 IEMOEN	Applies to all cohorts below	None		Has not exhausted SOC
	Dose escalation - monotherapy			
	Dose expansion - monotherapy			

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

Trial	Cohort	Molecul	Ineligibility reasons
		ar	
METC 02 KAYRAS	Applies to all cohorts below	KRAS G12D	PD-L1 expression below minimum of 50.0
	Dose expansion - monotherapy - Colorectum		
	Dose expansion - monotherapy - NSCLC		No lung non-small cell carcinoma

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

Cohort Trial Molecula Sites Configuration

None