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Radiomics Response Signature for Identification of Metastatic Colorectal Cancer Sensitive to Therapies Targeting EGFR Pathway

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Abstract

Background: The authors sought to forecast survival and enhance treatment decisions for patients with liver metastatic colorectal cancer by using on-treatment radiomics signature to predict tumor sensitiveness to irinotecan, 5-fluorouracil, and leucovorin (FOLFIRI) alone (F) or in combination with cetuximab (FC). Methods: We retrospectively analyzed 667 metastatic colorectal cancer patients treated with F or FC. Computed tomography quality was classified as high (HQ) or standard (SD). Four datasets were created using the nomenclature (treatment) – (quality). Patients were randomly assigned (2:1) to training or validation sets: FCHQ: 78:38, FCSD: 124:62, FHQ: 78:51, FSD: 158:78. Four tumor-imaging biomarkers measured quantitative radiomics changes between standard of care computed tomography scans at baseline and 8 weeks. Using machine learning, the performance of the signature to classify tumors as treatment sensitive or treatment insensitive was trained and validated using receiver operating characteristic (ROC) curves. Hazard ratio and Cox regression models evaluated association with overall survival (OS). Results: The signature (area under the ROC curve [95% confidence interval (CI)]) used temporal decrease in tumor spatial heterogeneity plus boundary infiltration to successfully predict sensitivity to antiepidermal growth factor receptor therapy (FCHQ: 0.80 [95% CI = 0.69 to 0.94], FCSD: 0.72 [95% CI = 0.59 to 0.83]) but failed with chemotherapy (FHQ: 0.59 [95% CI = 0.44 to 0.72], FSD: 0.55 [95% CI = 0.43 to 0.66]). In cetuximab-containing sets, radiomics signature outperformed existing biomarkers (KRAS-mutational status, and tumor shrinkage by RECIST 1.1) for detection of treatment sensitivity and was strongly associated with OS (two-sided P < .005). Conclusions: Radiomics response signature can serve as an intermediate surrogate marker of OS. The signature outperformed known biomarkers in providing an early prediction of treatment sensitivity and could be used to guide cetuximab treatment continuation decisions.

Colorectal cancer (CRC) is a leading cause of cancer death globally. Liver metastasis affects more than one-half of CRC patients (1). Anti-epidermal growth factor receptor (EGFR) therapies, including tyrosine kinase inhibitors and monoclonal antibodies, demonstrate activity in both CRC and other tumor types. In metastatic CRC (mCRC) patients, the assessment of anti-EGFR monoclonal antibody efficacy relies on computed tomography (CT) scan response endpoints. The on-treatment shrinkage of metastases on CT scans is considered a hallmark of EGFR signaling pathway dependency (2–5) and of treatment sensitivity (6). In unresectable mCRC, tumor shrinkage guides the clinical decision to pursue a curative opportunity (downstage to

resection) or palliative treatment to maximize overall survival (OS) and improve quality of life by symptom relief (1). As the decision to continue EGFR-targeted therapy must balance the risks and potential rewards of treatment, there is a pressing need for biomarkers that can estimate the likelihood of clinical benefit in individual patients.

We sought to meet this need for alternative biomarkers by using artificial intelligence (AI). We utilized machine learning to create an AI signature that evaluated a change in tumor phenotype between baseline and 8 weeks on CT-scan images to predict clinical outcome (OS). Radiology is undergoing rapid change because of remarkable advances in the field of AI, particularly

algorithms for machine learning and deep learning, which enable automated, high-throughput quantification of medical images as a set of quantitative features. Traditional radiomic features are identified according to a priori definitions (eg, tumor diameter or density), and artificial neural networks adaptively define deep learning features as spatial hierarchies of representations. The dataset of radiomic and deep learning features can be mined by machine learning to identify statistically significant associations between variables of interest, such as clinical outcome or tumor mutational status. This big data approach is applicable to any imaging modality but requires large and consistent datasets, which CT scanning currently best provides because of its use as a standard of care in oncology. In particular, the widespread adoption of CT measurement of tumor diameter (eg, RECIST 1.1) as an endpoint in clinical trials has generated an invaluable resource for AI research. The current study leverages this resource by analyzing data from the multicenter clinical trial NCT00154102.

AI techniques allow objective and reproducible analysis of CT image characteristics, including imaging features not apparent to the human eye. The overall goal of this study is to determine whether AI techniques can offer oncologists additional clinical information regarding response assessment. Our test case is the decision to continue anti-EGFR treatment. We utilized data from mCRC patients treated with cetuximab in combination with FOLFIRI (irinotecan, 5-fluorouracil, and leucovorin) in the randomized multicenter CRYSTAL trial (NCT00154102). Data from this trial were used to obtain regulatory approval for the clinical use of cetuximab in mCRC. Pretreatment selection is currently based on a biopsy sample (primary tumor or metastasis) or on liguid biopsy. In approximately one-half of mCRCs patients (7), a RAS wild-type mutational status (KRAS and NRAS exons 2, 3, 4) guides the clinical use of cetuximab in addition to either FOLFIRI (8-11) or FOLFOX (12). This treatment has been found to prolong median OS by 8 months in RAS wild type (10) and increase overall response rate, but also to increase the rate of adverse events (8). However, several tumor biopsy-driven clinical trials have shown the limitations of predicting clinical benefit from EGFR-targeted therapies in mCRC and other solid tumors based on a single biomarker (13). The exception is the presence of EGFR mutation in non-small cell lung cancer, which does predict a higher response rate and prolonged survival (13).

To determine whether an AI-driven signature could guide clinicians to continue EGFR-targeted therapies on an individualized basis, we aimed to develop and validate an on-treatment signature detecting mCRC patients sensitive FOLFIRI+cetuximab (FC) using quantitative assessment of tumor changes between baseline and 8-week CT images.

Methods

Study Design

Our primary endpoint was to train and validate an ontreatment signature detecting mCRC patients sensitive to FC. Our secondary endpoint was to generalize the on-treatment signature to other datasets with different quality of imaging settings and treatment types (Figures 1 and 2).

Participants and Datasets

We retrospectively analyzed one clinical trial involving two treatment arms and 667 CRC patients. The two treatment arms

involved anti-EGFR treatment regimens (cetuximab: C) and FOLFIRI (F). The CRYSTAL trial (NCT00154102) was a randomized, open-label, multicenter phase III trial comparing FC (cohort FC) with FOLFIRI alone (cohort F) in the first-line treatment of mCRC (Table 2). Data were gathered up to the clinical trials completion date. Patients with missing data were excluded. More information on this clinical trial can be found in the Supplementary Methods (available online).

We have evaluated the quality of CT-scan acquisition. More information about CT-scan characteristics, CT-scan quality (14-16), and radiomics quality score (17) can be found in Supplementary Methods (available online). Our goal was to create the signature in a dataset with high signal-to-noise ratio because of high imaging quality (HQ dataset) and evaluate if it can be more universally applicable to a dataset with standard qual-

Using the nomenclature (tumor type) – (treatment) – (quality of the dataset), the two cohorts (CRC-FC, CRC-F) were divided into four datasets and patients (patients) were randomly assigned to training and validation sets using a 2:1 ratio (CRC-FCHQ: 78:38 patients, CRC-FCSD: 124:62 patients, CRC-FHQ: 78:51 patients, CRC-F^{SD}: 158:78 patients) (Table 1; Figure 1).

Reference Standard: Sensitivity vs Resistance to **Systemic Cancer Therapy**

The signature was trained and validated to predict tumor sensitivity to systemic anticancer treatment. All cancer patients were divided into two groups: sensitive to treatment and not sensitive (resistant) to treatment (Supplementary Methods, available online).

In mCRC patients (datasets CRC-FCHQ, CRC-FCSD, CRC-FHQ, CRC-F^{SD}), the reference standard to determine tumor sensitivity to treatment was OS. Patients were considered resistant if they died before 17.7 months (median OS in the training set) and were considered sensitive otherwise. The failure event for OS was defined as death due to any cause. Survival time was measured from the date of random assignment to the date of death or last follow-up (censored observation). No included patients were censored before 17.7 months: all patients were followed beyond 17.7 months unless they died due to any cause.

Predictors Used in the Signature: Radiomics and Deep-**Learning Features**

In total, 3499 quantitative image features including 1757 radiomics features and 1742 pretrained deep-learning features were used to characterize the early CT changes in tumor phenotype. The full details of lesion segmentation and feature extractions can be found in the Supplementary Methods (available online).

To reduce overfitting, a radiomics signature integrated up to four of these features. These four were those ranked highest by the machine learning algorithm in terms of prediction importance (18) using the best predictive model developed in the training set of the discovery cohort CRC-FC $^{\rm HQ}$. The full details of model building can be found in the Supplementary Methods (available online).

Signature Building in Cohort CRC-FCHQ

In the training set of the discovery cohort (CRC-FC^{HQ}), we developed a multivariable prediction model (ie, the signature, to

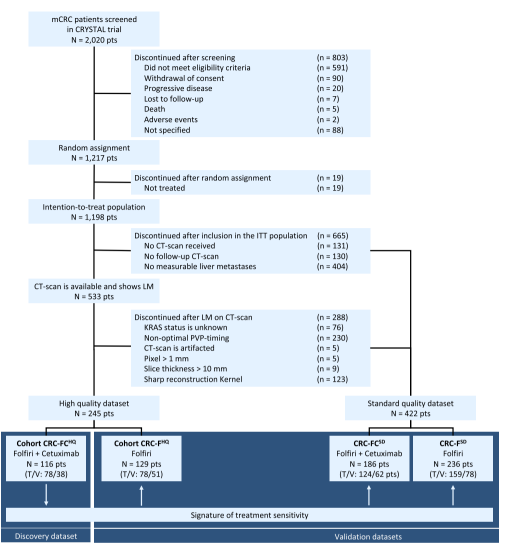


Figure 1. CONSORT diagram. Patients could be excluded for multiple reasons. The withdrawal boxes show the number of patients excluded at each step. LM = liver metastasis; T = training set; V = validation set. The artificial intelligence (AI) signature was developed (training set) and validated (validation set) in the FOLFIRI+cetuximab discovery cohort with high-quality (HQ) dataset.

predict tumor sensitivity to treatment) (see Figure 2). In the implementation, a "coarse" to "fine" strategy was developed to select optimal features from the large number of extracted quantitative image features to build the signature. The coarse selection approach consists of three processes (reproducibility analysis, redundancy analysis, and feature ranking) aiming to screen many nonreproducible, redundant, and noninformative candidate features. The fine selection approach consists of a "forward" search and feature combination, aiming to select optimal features and build the final optimal model.

Signature Calibration in Cohorts CRC-FCSD, CRC-FHQ, and CRC- F^{SD}

The four imaging biomarkers identified in the signature developed in the training set of CRC-FC^{HQ} were computed in the training sets of the cohorts CRC-FC^{SD}, CRC-F^{HQ}, and CRC-F^{SD}. The signature developed in the training set of $CRC\text{-}FC^{HQ}$ was transferred and calibrated to predict tumor sensitivity to treatment in

the training sets of the cohorts CRC-FCSD, CRC-FHQ, and CRC-FSD as described in Supplementary Methods (available online).

Validation of the Signature

The primary endpoint was to evaluate the performance of the signature to predict tumor sensitivity to treatment. The performance was measured using area under the receiver operating characteristic curve (AUC) in the validation sets of the four cohorts consisting of patients that were not used for training.

The methodology used for secondary endpoints is described in Supplementary Methods I.6.1 (available online).

Statistical Analysis

Statistical analysis was conducted using Matlab (version 2016b; Mathworks, Natick, MA) and SPSS (version 23.0; IBM; Armonk, NY). AUC and AUC's 95% confidence interval (CI)

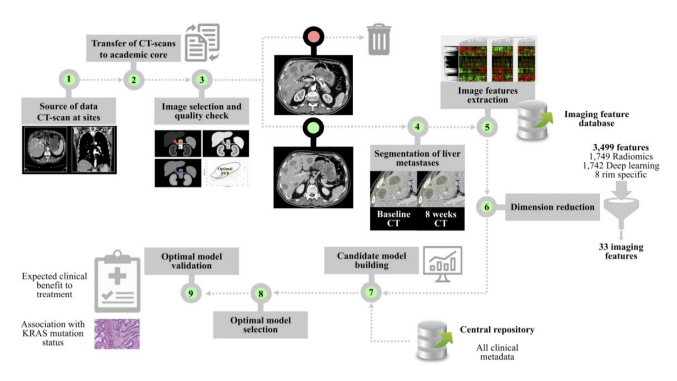


Figure 2. Artificial intelligence (AI) workflow. Steps 1-2: Computed tomography (CT) scans acquired at study sites are transferred to our academic core. Step 3. Image selection and quality check using a computer-aided algorithm designed by machine-learning. Step 4. Segmentation of liver metastases on CT scan by an expert radiologist at baseline and 8 weeks in each patient. Step 5. Combination of all segmented lesions to compute a tumor imaging phenotype in each patient based on imaging features extraction in each segmented liver metastasis (3499 imaging features characterizing changes between baseline and 8 weeks). Step 6. Dimension reduction using machine learning. Identification of reproducible, nonredundant and informative candidate imaging features for model building. Step 7. Candidate model building to enhance strategic decision-making (training set). Step 8. Optimal model selection in the training set using threefold cross-validation to evaluate the performance of candidate models in terms of area under the receiver operating characteristic curve. Step 9. Signature validation (validation set).

Table 1. Patient characteristics and cohorts in the CRYSTAL trial (NCT00154102)

Treatment arm	Cohort FOLFIRI plus cetuximab	Cohort FOLFIRI alone	
No.	Dataset CRC-FC ^{HQ} : 116 patients Dataset CRC-FC ^{SD} : 186 patients	Dataset CRC-F ^{HQ} : 129 patients Dataset CRC-F ^{SD} : 236 patients	
Tumor characteristics	•	•	
Tumor type	CRC	CRC	
Stage	Advanced	Advanced	
Lesion(s) segmented	Liver metastases	Liver metastases	
Clinical trial characteristics			
Biomarker	KRAS* WT mutational status	KRAS* WT mutational status	
Treatment regimen			
Chemotherapy	FOLFIRI	FOLFIRI	
Anti-EGFR	Cetuximab (anti-EGFR mAb)	None	
Primary endpoint			
Outcome	OS, PFS	OS, PFS	
Sensitivity to anti-EGFR mAb	OS benefit compared with FOLFIRI alone	N/A	
Time of completion of CT scans	Baseline and 8 weeks	Baseline or 8 weeks	
Clinical trial no.	NCT00154102	NCT00154102	

^{*}KRAS status and not RAS status was evaluated post hoc in the CRYSTAL clinical trial. CRC = colorectal cancer; F = FOLFIRI; FC = FOLFIRI+cetuximab; HQ = High com $puted\ tomography\ quality;\ SD = Standard\ computed\ tomography\ quality;\ mAb = monoclonal\ antibody.$

were used to indicate classification performance. Analysis of variance was performed to compare continuous variables, and χ^2 test was performed to compare categorical variables. Cox regression was used to investigate the effect of survival variables, and log-rank test was used to compare survival times of two groups. A P less than .05 was used to determine statistical significance. All tests were two-sided.

Results

Participants

Table 2 shows that no statistically significant differences were detected by the post hoc exploratory tests performed between cohorts CRC-FCHQ, CRC-FCSD, CRC-FHQ, and CRC-FSD and

Table 2. Patient characteristics in the CRYSTAL trial (NCT00154102)*

	LM mCRC patients with liver metastases				SD quality
	HQ quality				
	Cohort C	Cohort CRC-FC ^{HQ} -FC		Cohort CRC-F ^{HQ} FOLFIRI	
Patient characteristics	Training No. (%)	Validation No. (%)	Training No. (%)	Validation No. (%)	and CRC-F No. (%)
Patients, no.	78	38	78	51	288
Gender					
Male	50 (64.1)	29 (76.3)	51 (64.1)	31 (60.8)	184 (63.9)
Female	28 (35.9)	9 (23.7)	27 (34.6)	20 (39.2)	104 (36.1)
Primary site	()	- ()	(/	. (/	(****)
Right colon	20 (25.6)	14 (36.8)	20 (25.6)	17 (33.3)	70 (24.3)
Left colon	58 (74.4)	24 (63.2)	58 (74.4)	34 (66.7)	217 (75.3)
Prior adjuvant chemotherapy	38 (74.4)	24 (03.2)	36 (74.4)	3 1 (00.7)	217 (73.3)
Yes	0 /11 E\	C (1E 0)	10 (12 0)	<i>C (</i> 11 0)	40 (12 0)
	9 (11.5)	6 (15.8)	10 (12.8)	6 (11.8)	40 (13.9)
No	69 (88.5)	32 (84.2)	68 (87.2)	45 (88.2)	248 (86.1)
Leucocytes	CE (00.0)	05 (00.4)	CE (00.0)	04 (66 7)	004 (04.0)
Low	65 (83.3)	35 (92.1)	65 (83.3)	34 (66.7)	234 (81.3)
High LDH baseline	12 (15.4)	3 (7.9)	10 (12.8)	15 (29.4)	45 (15.6)
Low	41 (52.6)	18 (47.4)	28 (35.9)	21 (41.2)	147 (51.0)
High	29 (37.2)	15 (39.5)	36 (46.2)	21 (41.2)	111 (38.5)
Unknown	8 (10.3)	5 (13.2)	14 (17.9)	9 (17.6)	30 (10.4)
Maximum staining intensity	` '	` '	, ,	, ,	, ,
Low	13 (16.7)	1 (2.6)	12 (15.4)	7 (13.7)	48 (16.7)
High	65 (83.3)	37 (97.4)	65 (83.3)	44 (86.3)	238 (82.6)
Histology of primary tumor	()	()	()	()	
Adenocarcinoma	72 (92.3)	35 (92.1)	76 (97.4)	50 (98.0)	283 (98.3)
Mucinous adenocarcinoma	6 (7.7)	3 (7.9)	2 (2.6)	1 (2.0)	5 (1.7)
Histological differentiation	0 (7.7)	3 (7.5)	2 (2.0)	1 (2.0)	3 (1.7)
Well differentiated	10 (12.8)	4 (10.5)	15 (19.2)	7 (13.7)	22 (11 5)
Moderately differentiated	, ,	, ,	, ,	, ,	33 (11.5)
•	51 (65.4)	24 (63.2)	50 (64.1)	33 (64.7)	191 (66.3)
Poorly differentiated	15 (19.2)	6 (15.8)	12 (15.4)	8 (15.7)	44 (15.3)
KRAS BRAF mutation status	05 (00.4)	10 (50 0)	0.5 (0.0.0)	47 (00.0)	75 (05 0)
KRAS mutant	25 (32.1)	19 (50.0)	26 (33.3)	17 (33.3)	75 (26.0)
KRAS and BRAF WT	50 (64.1)	15 (39.5)	46 (59.0)	31 (60.8)	129 (44.8)
BRAF mutant	3 (3.8)	4 (10.5)	6 (7.7)	3 (5.9)	8 (2.8)
Unknown	0 (0.0)	0 (0.0)	0 (0.0)	0 (0.0)	76 (26.4)
Target lesions at baseline, no.					
0	0 (0.0)	1 (2.6)	0 (0.0)	1 (2.0)	2 (0.7)
1	5 (6.4)	5 (13.2)	12 (15.4)	3 (5.9)	31 (10.8)
2	9 (11.5)	10 (26.3)	13 (16.7)	11 (21.6)	45 (15.6)
3	14 (17.9)	7 (18.5)	18 (23.1)	11 (21.6)	44 (15.3)
4	13 (16.7)	4 (10.5)	5 (6.4)	6 (11.8)	43 (14.9)
5	24 (30.8)	7 (18.5)	16 (20.5)	10 (19.6)	70 (24.3)
>5	14 (17.9)	4 (10.5)	14 (17.9)	10 (19.6)	53 (18.4)
Nontarget lesions at baseline, no.					
0	17 (21.8)	16 (42.1)	20 (25.6)	16 (31.4)	77 (26.7)
1	35 (44.9)	8 (21.1)	25 (32.1)	16 (31.4)	90 (31.3)
2	10 (12.8)	4 (10.5)	22 (28.2)	11 (21.6)	50 (17.4)
3	6 (7.7)	6 (15.8)	6 (7.7)	4 (7.8)	19 (6.6)
4	2 (2.6)	1 (2.6)	2 (2.6)	1 (2.0)	13 (4.5)
5	4 (5.1)	2 (5.3)	2 (2.6)	1 (2.0)	18 (6.3)
>5	4 (5.1)	1 (2.6)	1 (1.3)	2 (3.9)	21 (7.3)
Only liver metastasis	57 (73.1)	27 (71.1)	49 (62.8)	37 (72.5)	21 (7.3)
-					· ·
Only liver and lung metastasis	63 (80.8)	33 (86.8)	73 (93.6)	45 (88.2)	253 (87.8)

Liver metastatic patients included in the HQ cohorts have similar characteristics. The only difference is the quality of the CT scan acquisition protocol and reconstruction settings. The distribution of clinical characteristics for each subgroup of patients was compared using analysis of variance and χ^2 (P values) and did not demonstrate the distribution of clinical characteristics for each subgroup of patients was compared using analysis of variance and χ^2 (P values) and did not demonstrate the distribution of clinical characteristics for each subgroup of patients was compared using analysis of variance and χ^2 (P values) and did not demonstrate the distribution of clinical characteristics for each subgroup of patients was compared using analysis of variance and χ^2 (P values) and did not demonstrate the distribution of clinical characteristics for each subgroup of patients was compared using analysis of variance and χ^2 (P values) and did not demonstrate the distribution of clinical characteristics for each subgroup of patients was compared using an algorithm. $strate\ any\ statistically\ significant\ difference\ between\ the\ cohorts.\ CT=computed\ tomography;\ CRG=colorectal\ cancer;\ F=FOLFIRI;\ FC=FOLFIRI+cetuximab;\ HQ=folderset for the cohorts.$ $\label{eq:bound} \mbox{High computed tomography quality; SD} = \mbox{Standard computed tomography quality.}$

Table 3. Performance of the AI signature in the validation sets*

	Anti-EGFR treatment		No anti-EGFR treatment		
	FOLFIRI +	cetuximab	FOLFIRI only		
	mCRC		mCRC		
Analysis	CRC-FC ^{HQ}	CRC-FC ^{SD}	CRC-F ^{HQ}	CRC-F ^{SD}	
Reference standard	OS	OS	OS	OS	
No. patients					
Total	116	186	129	236	
Training	78	124	78	159	
Sensitive	48	71	42	85	
Resistant	30	53	36	74	
Validation	38	62	51	78	
Sensitive	23	34	32	42	
Resistant	15	28	19	36	
Signature					
Algorithm	Random Forest	Random Forest	Random Forest	Random Forest	
Features	Shape SI4	Shape SI4	Shape SI4	Shape SI4	
	LoG Z Entropy	LoG Z Entropy	LoG Z Entropy	LoG Z Entropy	
	GTDM Contrast	GTDM Contrast	GTDM Contrast	GTDM Contrast	
	LoG X Entropy	LoG X Entropy	LoG X Entropy	LoG X Entropy	
Performance					
Tumor sensitivity	OS >17.7 mo	OS >17.7 mo	OS >17.7 mo	OS >17.7 mo	
AUC, training	0.83 (95% CI = 0.75 to 0.95)	0.84 (95% CI = 0.76 to 0.89)	0.75 (95% CI = 0.63 to 0.85)	0.75 (95% CI = 0.67 to 0.82)	
AUC, validation	0.80 (95% CI = 0.69 to 0.94)	0.72 (95% CI = 0.59 to 0.83)	0.59 (95% CI = 0.44 to 0.72)	0.55 (95% CI = 0.43 to 0.66)	
Association with OS Cox regression					
Hazard ratio	44.3 (95% CI = 6.4 to 307.7)	6.5 (95% CI = 1.8 to 23.6)	1.9 (95% CI = 0.4 to 8.4)	0.96 (95% CI = 0.2 to 4.3)	
Death	31/38	53/62	42/51	69/78	
Р	.0001	.005	.43	.96	

^{*}The performance of the AI signature to detect treatment sensitivity was evaluated in all cohorts. AI = Artificial Intelligence; AUC = Area Under the Curve; CT = computed tomography; CRC = colorectal cancer; F = FOLFIRI; FC = FOLFIRI+cetuximab; HQ = High computed tomography quality; SD = Standard computed tomography quality.

between training and validation sets for each treatment arm. Supplementary Figures 2 and 3 (available online) show tumor imaging-phenotyping using unsupervised analysis. Supplementary Table 3 (available online) shows that features were different between HQ and SD datasets; hence there was a need for a calibration in each cohort.

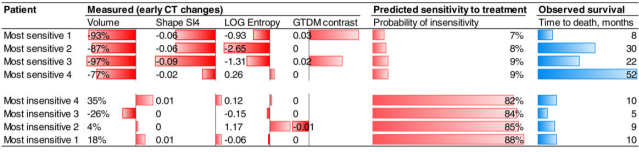
Performance of the Signature in Cohort CRC-FC^{HQ}

Of 116 patients, 95 patients died during the follow-up. The median follow-up duration was 21.8 months overall and 46.7 months (range = 22.4-55.0) in censored patients. All patients with censored survival data had follow-up longer than 17.7 months and belonged to the low-risk group.

In the training set, the best predictive model was built using a Random Forest algorithm (18) and 4 Radiomics features (Figures 3). The performance of the signature to diagnose treatment-sensitive tumors was AUC = 0.83 (95% CI = 0.75 to 0.92); sensitivity = 0.77, and specificity = 0.85. 4 Radiomics features were included in the signature (Table 3; Figures 3): Shape SI4, LoG Z Entropy, LoG X Entropy, and GTDM Contrast. These four features were transferred to all cohorts. Shape SI4 characterizes the local curvatures in surface shape of a three-dimensional object. Entropy is a metric of the disorder or heterogeneity of the signal that is computed after the application of a Gaussian smoothing filter to reduce noise and a Laplacian operator to highlight regions of rapid gray-level change in images. Gray Tone Difference Matrix features describe visual properties of texture based on gray-tone difference between a pixel and its neighborhood. Definitions are presented in Supplementary Methods (available online).

In the validation set, the performance of the signature (AUC = 0.80, 95% CI = 0.69 to 0.94, sensitivity = 0.80, and specificity =0.78) statistically significantly (two-sided P < .001) outperformed both KRAS mutational status at baseline (AUC = 0.67, P < .001) and 8-week tumor shrinkage (AUC = 0.75 using either RECIST 1.1-like unidimensional tumor shrinkage or volumetric tumor shrinkage) (P < .001 using bootstrap; Supplementary Figure 6, available online). There was no biological or clinical phenotype of failure (Supplementary Table 2, available online).

In the cohort CRC-FC^{HQ}, the signature showed good prognostic discrimination for OS (P < .001) by univariate Cox regression analysis. High-risk signature (signature > 0.5) was associated with shorter OS than patients with low-risk signature in the training set, the validation set, KRAS wild-type patients, and KRAS mutant tumors (Figure 4) (P < .001). The signature measured at 8 weeks in KRAS wild-type patients can be used to predict survival at any time (nomogram, Supplementary Figure 7, available online). The signature provided incremental value over existing benchmarks for the prediction of OS because univariate Cox regression analyses adjusted for KRAS mutational status demonstrated that the signature had a statistically significantly better fit for the prediction of



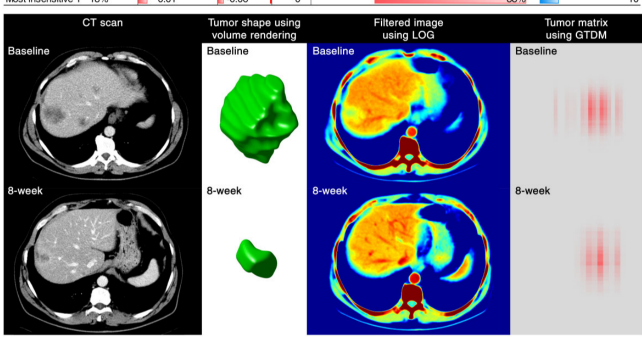


Figure 3. Visual representation of the four imaging features included in the signature. The changes in the radiomics features in the patients with the lowest and the highest probability of insensitivity to treatment according to the radiomics signature are presented on this graph (most sensitive 1-4 vs most insensitive 1-4). The changes in tumor imaging phenotype of the patient "most sensitive 1" is displayed below. As demonstrated, CT-scan images are transformed to other mathematical spaces for feature extraction, for example, CT image is transformed to LOG space for computing the entropy value (spatial heterogeneity), and tumor pixels within segmentation contour are transformed to GTDM matrix for computing the contrast value

patients' OS than tumor shrinkage (P < .001) (Supplementary Figure 6; Supplementary Tables 1, 3, and 4, available online).

Generalizability of the Signature to CRC-FC $^{\rm SD}$, CRC-F $^{\rm HQ}$. and CRC-F^{SD}

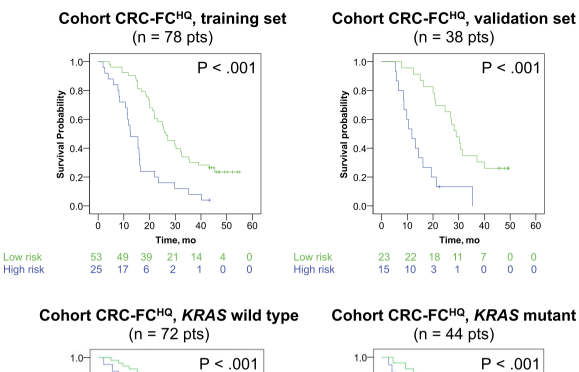
The Random Forest algorithm (18) using the 4 Radiomics features signature developed and used in the $\mathsf{CRC}\text{-}\mathsf{FC}^\mathsf{HQ}$ analysis (Supplementary Figure 4, available online) was applied to the other training datasets for calibration. The performance of the signature (Table 3; Figure 5; Supplementary Figure 8; Supplementary Tables 5 and 6, available online) in the validation sets consisting of unseen data was calculated. AUC was 0.72 (95% CI = 0.59 to 0.83), sensitivity = 0.82, and specificity = 0.61 in CRC- FC^{SD} . AUC was 0.59 (95% CI = 0.44 to 0.72), sensitivity = 0.37, and specificity = 0.84 in CRC-F^{HQ}. AUC was 0.55 (95% CI = 0.43 to 0.66), sensitivity = 0.81, and specificity = 0.33 in CRC-FSD.

Discussion

Using standard-of-care CT scans acquired in multicenter clinical trials, AI successfully identified on-treatment imaging

biomarkers associated with tumor sensitivity to anti-EGFR therapy in CRC patients. These four imaging biomarkers enabled development of a signature, which offered high sensitivity and specificity, to guide the clinical decision to continue EGFR targeted therapies. The innovation of our work compared with the existing literature was that the signature was dynamic, generalized to four datasets acquired in multicenter studies, used a limited subset of features to reduce overfitting, and considered image quality according to new quality control tools (17,19,20). Because it is based on routinely acquired CT scans, such a tool could be widely incorporated into clinical practice at minimal cost to detect EGFR-resistant tumors due to clonal acquisition of resistance mechanisms through alternate pathways and downstream events.

The signature features can be understood as noninvasive in vivo surrogates of treatment-induced biological changes and generally fall into two categories: indicators of heterogeneity of the metastatic burden and evaluation of tumor-parenchyma interactions. In patients with both KRAS wild-type or mutated tumors, the imaging features identified by our signature provided incremental value over 8-week tumor shrinkage (6) in a multifactorial prediction model. Although KRAS wild-type status at baseline was associated with better outcome to



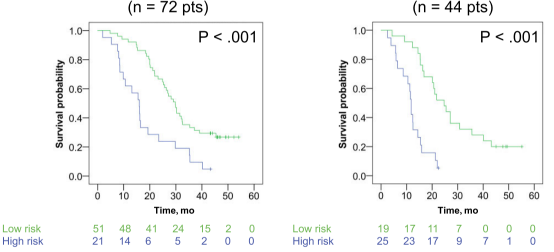


Figure 4. Risk stratification using the signature in the discovery cohort. The discovery cohort included mCRC patients treated with FOLFIRI+ cetuximab. Kaplan-Meier graphs depicting overall survival in patients stratified at high-risk (signature >0.5) or low-risk (signature ≤0.5) at 8 weeks by the signature. P values are based on data-set-stratified two-sided log-rank tests. A, B) Survival probability in the training and in the validation sets. C, D) Survival probability in KRAS wild-type and KRAS mutant groups. CRC = colorectal cancer; F = FOLFIRI; FC = FOLFIRI+cetuximab; HQ= High computed tomography quality; SD = Standard computed tomography quality.

treatment, adding this mutational status to the signature did not contribute to the incremental value. The capacity of the complete RAS mutation status to improve predictability was not assessed in our research. Although this finding is consistent with the literature suggesting that on-treatment markers of efficacy are better predictors of outcome than baseline variables, it also indicates that imaging biomarkers can overcome two weaknesses associated with the use of a single genomic biomarker to predict outcome to targeted therapies: tumor heterogeneity and pathways complexity.

Because CT scans can provide information regarding the entire tumor burden and be repeated noninvasively, they are better suited to address the spatial and temporal heterogeneity of tumors than genomic analysis of tissue from a single biopsy site. First, although KRAS mutations are early driving events in colorectal tumorigenesis and progression, substantial spatial heterogeneity exists in KRAS status. Despite good concordance between the primary tumors and matched distant metastases,

KRAS status shows 8% intratumoral heterogeneity in the primary tumor, up to 10% variability between the primary tumor and metastases, and up to 12% heterogeneity among multiple metastatic lesions (21). Second, the limitations of genomic biomarkers are further demonstrated by the fact that failure of anti-EGFR therapy is observed in one-half of the patients with wild-type RAS CRC. Although downstream mutations in genes (eg, BRAF and PI3KCA mutations) are not predictive biomarkers for anti-EGFR therapy, they seem to predict a poor prognosis independent of treatment. Third, temporal heterogeneity may also emerge from the inherent dynamic evolution and adaptation of clones acquiring distinct resistance mechanisms, especially in the presence of selection pressure from anti-EGFR targeted therapies (22).

Decrease in heterogeneity of the metastatic burden (eg, entropy) on CT scan following cetuximab treatment was the best predictor of tumor sensitivity to anti-EGFR therapies. The signature can be understood as an AI analysis of CT image

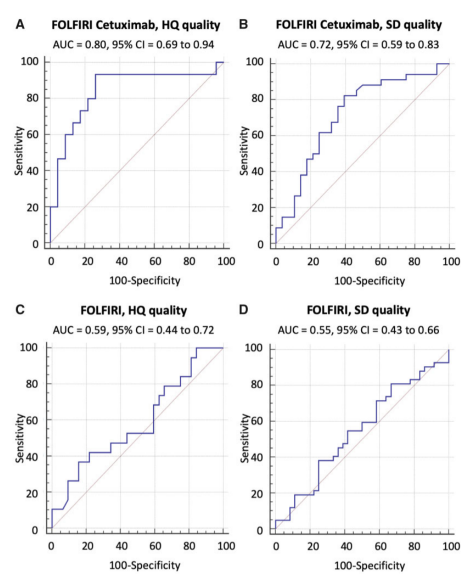


Figure 5. Performance of the signature in the validation sets of the three independent testing cohorts. CT quality was classified as high (HQ) or standard (SD). The Random Forest algorithm using the 4 Radiomics features signature developed and used in the FCHQ analysis was applied to the other training data sets for calibration. Then, its performance was validated in the four validation sets. Area under the receiver operating characteristic curve (AUC) and AUC's 95% confidence interval (CI) were used to indicate classification performance. The signature (AUC [95CI]) used temporal decrease in tumor spatial heterogeneity plus boundary infiltration to successfully predict sensitivity to anti-EGFR therapy (FCHQ [A], FCSD [B]) but failed with chemotherapy (FHQ [C], FSD [D]). F = FOLFIRI; FC = FOLFIRI+cetuximab; HQ= High computed tomography quality; SD = Standard computed tomography quality.

heterogeneity (23) capturing a key hallmark of cancer (24): the heterogeneity of an expanding aberrant and distorted neovasculature supplying tumor growth. These macroscopic patterns of neovascularization are marked by heterogeneous and excessive blood flow, microhemorrhage, and leakiness, which produce a heterogeneous accumulation of iodine on contrastenhanced CT scans. Radiomics signature can therefore capture neovascularization patterns (14,25-27) associated with reduced treatment efficacy (25,28), poor outcome because of reduced drug delivery (29-32), hypoxia, and promotion of immune evasion, tumor progression, and metastasis (24). From a broader perspective, there is a rationale that even basic CT features such as size changes are associated with pathologic response (ie, the percentages of residual tumor) and OS in mCRC patients treated with chemotherapy regimens containing molecular targeted therapies targeting such as anti-EGFR or anti-VEGF (33).

The second weakness of the single genomic biomarker is the complexity of cancer pathways. Following blockade of oncogenic pathways by targeted therapies, the Darwinian selection process will tend to select cancer cells that exploit alternative pathways. AI analysis of CT scans can identify signatures associated with downstream events linked to favorable outcomes with anti-EGFR therapy. Several tumor biopsy-driven clinical trials (13) demonstrated that inhibition of the EGF receptor activation alone did not confirm clinical benefit in mCRC and other solid tumors. This is explained by the selection of clones with downstream mutations allowing the tumor to continue to proliferate (eg, EGFR, then RAS, then RAF or PI3K). A radiomics signature can dynamically assess features associated with efficient blockade of the full range of oncogenetic pathways, and their interconnections promise to accelerate the development of molecularly targeted therapies.

Tumor shape (eg, shape SI4) and boundary sharpness (eg, Sigmoid Slope Energy) features were strongly associated with OS and capture complex patterns of macroscopic changes at tumor-liver interfaces in mCRC that reflect both infiltrative tumor growth and perfusion change. A pushing interface in which tumor cells directly compress the liver has a poorer prognosis than interfaces in which a desmoplastic lymphocytic stroma separates the tumor cells from the liver or in which tumor cells replace liver parenchyma without pushing or desmoplastic reaction (34). Conversely, the presence of a peritumoral contrast-enhancement rim region surrounding liver tumors is biologically relevant as a reflection of recruitment of hepatic artery branches by vascular endothelial growth factor and/or a reduced peritumoral portal flow (mechanical tumor compression and/or vessel lumen-narrowing secondary to fixed leukocyte adherence) (16,35-40).

There are several innovative aspects of our work. Although AI techniques are increasingly proposed to aid drug discovery and clinical imaging, none have yet been identified and robustly validated in heterogeneous datasets such as large multicenter clinical trials. Our innovative methodology (41–43) lies in combining the deep-learning feature set (ImageNet pretrained VGG-16) and radiomics feature set (engineered features) and applying them using a dynamic approach (two time points) to response assessment in cancer therapies. Our results suggest that the performance of imaging-based response assessment could progressively increase by moving from unidimensional size, to volume (44), and finally by integrating radiomics features. Another innovation—compared with other radiomic signatures in CRC patients (45-47)—is that we tightly controlled the quality (17) of image acquisition through robust reconstruction settings (15), reproducible segmentation (16), and selection of optimal contrast enhancement at the portal venous phase (14). This was by design to increase the signal-to-noise ratio because of the heterogeneity in acquisition protocols and image quality in a landmark historical multicenter clinical trial with "real-life" heterogeneous image quality. Finally, we reduced the risks of type I error and overfitting by using only four imaging biomarkers in the signature and by evaluating the generalizability of these imaging biomarkers to four validation sets.

This work has limitations but opens numerous perspectives and opportunities for improvement. First, future research could assess the performance of CT scan evaluation earlier than 8 weeks or pool data from multiple timepoints and modeling the growth and decay of radiomics features for all available patient scans beyond week 8. Second, before prospectively implementing this decision tool in RAS wild-type patients, we expect to calibrate the signature to all generations of CT scanners and fine-tune convolutional neural network models in external and larger datasets (48). Third, we will evaluate the possible temporal within-patient heterogeneity of responses to identify therapy-resistant clones. Fourth, there is a need to determine if these imaging biomarkers uncover a prognostic effect rather than predictive (to anti-EGFR therapy), specifically in certain subpopulations. As an ancillary study, AI could not identify a phenotype associated with the sensitivity of tumors in the FOLFIRI-only group in a clinically significant manner. Because CRC is known to evolve by a reiterative process of genetic diversification and clonal evolution, a targeted therapy efficient in only some patients might indeed produce greater interpatient variability in terms of depth of response than chemotherapy (49). This is explained by the fact that sensitive tumors will be more homogeneous than insensitive tumors because resistant mutation events will develop heterogeneous resistant tumor niches.

Temporal changes in the spatial heterogeneity of metastatic tumor burden were linked to outcome. Heterogeneity was defined by the conversion of a per-lesion phenotype to a perpatient phenotype. First, we calculated the spatial heterogeneity between neighboring voxels in each metastatic lesion. Then, we aggregated all metastatic lesions through a weighted average. Of note, each liver metastatic lesions may theoretically behave differently from metastatic disease in peritoneum and lungs. Nonetheless, several facts support our methodology. First, liver metastasis affects greater than one-half of CRC patients (1). Second, a modeling framework (22)—considering resistance development and clonal selection—demonstrated similar dynamics among individual lesions within the same tumor site location and different dynamics in only 35% of patients with lesions in different anatomic locations (22).

Although there was no apparent selection bias, the signature was designed for CRC patients with liver metastases with known KRAS status in a landmark trial (48–51). The architecture of the liver parenchyma differs from other tissues such as the lung parenchyma. Therefore, some imaging features (ie, shape and boundary sharpness features) might intuitively need to be calibrated for each malignant site, whereas surrogates of heterogeneity may generalize. We anticipate that features identified in this historical landmark clinical trial can be easily generalized and fine-tuned to analyze newer datasets. First, features were generalizable from the HQ cohort to patients with SD quality imaging settings. Second, response evaluation on CT scan has not fundamentally changed over the last decades since new technologies focused on increasing the speed and reducing radiation dose (50). The main advantage of using an "old" trial is the extended follow-up of patients that allowed only 18% of data to be censored for the OS. Most newer clinical trial datasets are not yet mature or available from sponsors; nevertheless, groups are actively attempting to make these publicly available (51). In conclusion, AI-derived support tools could give clinicians an early prediction of the success of treatment with FC using conventional standard-of-care CT scans.

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