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## Abstract

Compared to other malignancies, there is a lack of easy-to-evaluate biomarkers for gastric cancer, which is associated with an adverse clinical outcome in many cases. Here, we present Stroma AReactive Invasion Front Areas (SARIFA) as a new histological prognostic marker. We defined SARIFA as the direct contact between a cluster of tumor glands/cells comprising at least five tumor cells and inconspicuous surrounding adipose tissue at the invasion front. A total of 480 adenocarcinomas of the stomach and the gastroesophageal junction from two different collections were classified according to SARIFA. To understand the potential underlying mechanisms, a transcriptome analysis was conducted using digital spatial profiling (DSP). It was found that 20% of the tumors were SARIFA-positive. Kappa values between the three pathologists were good in both collections: 0.74 and 0.78. Patients who presented SARIFA-positive tumors had a significantly lower overall survival in Collections A (median: 20.0 versus 44.0 months; p = 0.014, n = 160) and B (median: 15.0 versus 41.0 months; p < 0.0001, n = 320). SARIFA positivity emerged as a negative independent prognostic factor for overall survival (HR 1.638, 95% Cl 1.153–2.326, p = 0.006). Using DSP, the most upregulated genes in SARIFA-positive cases were those associated with triglyceride catabolism and endogenous sterols. COL15A1, FABP2, and FABP4 were differentially expressed in positive cases. At the protein level, the expression of proteins related to lipid metabolism was confirmed. SARIFA combines low inter-observer variability, minimal effort, and high prognostic relevance, and is therefore an extremely promising biomarker related to tumor-promoting adipocytes in gastric cancer. © 2021 The Authors. The Journal of Pathology published by John Wiley & Sons, Ltd. on behalf of The Pathological Society of Great Britain and Ireland.

Keywords: gastric cancer; Stroma Areactive Invasion Front Areas; SARIFA; biomarker; histopathology; digital spatial profiling

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## Introduction

Gastric cancer is ranked as the sixth most common cancer entity worldwide, having accounted for approximately 780 000 cancer-associated deaths in 2018 [1]. So far, the best parameter for predicting prognosis, and therefore therapy, in gastric cancer patients is TNM staging. The factors that are relevant for determining the prognosis of gastric carcinomas are local infiltration depth, locoregional lymph node involvement, distant metastases, and vascular invasion [2–4]. Additionally, diffuse Laurén subtype and proximal tumor localization are also known negative prognostic factors for gastric cancer [5–7]. The introduction of perioperative chemotherapy after 2005 has improved the outcome in stage 2 and 3 gastric cancers, with a median survival of 50 months versus 34 months [8]. However, the prognosis of gastric cancer is still poor and has a 5-year survival rate that has not changed during the period between 2000 and 2014, with survival rates being between 31.4% and 33.5% in Germany [9]. To improve the prognosis estimation in gastric cancer beyond TNM staging, histomorphology-based concepts, such as tumor budding and the tumor–stroma ratio (TSR), have already been investigated. Unlike the case of colorectal cancer, such biomarkers have not reached general acceptance or been recommended in routine diagnostics [10–12]. Moreover, the molecular classification of the Cancer Genome Atlas (TCGA) identified distinct subgroups of

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© 2021 The Authors. The Journal of Pathology published by John Wiley & Sons, Ltd. on behalf of The Pathological Society of Great Britain and Ireland. This is an open access article under the terms of the Creative Commons Attribution-NonCommercial License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited and is not used for commercial purposes. gastric carcinomas that were associated with specific clinicopathological characteristics, but they failed to robustly predict patient survival [13]. So far, only microsatellite instability (MSI) and programmed death-ligand 1 (PD-L1) status have gained acceptance as prognostic biomarkers in gastric cancer [14]. Therefore, there is still a need to identify biomarkers for improving the prognostic prediction and therapy stratification in gastric cancer.

During our investigations addressing the prognostic value of the TSR in colon cancer, we recognized that in a certain portion of cases the tumor formations come into direct contact with adipose cells that lack a stromal reaction. Quite recently, we were able to confirm our hypothesis that this morphological feature, which we referred to as Stroma AReactive Invasion Front Areas (SARIFA), can serve as an adverse prognostic factor for the outcome in colon cancer [15]. The advantage of this biomarker is its simplicity; it does not need any additional testing or expenditure, offers ease of learning, has minimal time requirements, and provides extraordinarily low interobserver variability. The identification of this phenomenon in colon cancer gave rise to the question of whether this occurs in other cancer entities as well. The fact that SARIFA represents the areas in which there is direct contact between the tumor cells and the local mesenchymal tissue without intervening reactive collagen raises the question of whether this enables an interaction that has tumor-promoting effects. Considering the observation that obesity is associated with an increased incidence and aggressivity of malignant tumors, there is increasing interest in the crosstalk between the tumor cells and adipocytes. Indeed, the experimental data indicate that tumors gain advantages from a direct interaction with adipocytes [16].

The aim of the present study was to investigate the prognostic significance of SARIFA in a limited test collection of adenocarcinomas of the stomach and the gastroesophageal junction (n = 160) and to further validate the results in a larger collection of 320 patients. In addition, the possible underlying mechanisms or differences between negative and positive cases were investigated using transcriptome and subsequent immunohistochemical analyses.

## Materials and methods

#### Patients

Patient collection A served as the test collection and consisted of surgical resection specimens from 160 patients with adenocarcinomas of the stomach and the gastroesophageal junction (AEG II and III according to Siewert and Stein [17]) that at least presented infiltration of the submucosa (pT1b). These patients were treated between 2005 and 2018 at the Department of Surgery of the University Hospital Augsburg. In total, 69% of the tumors had been treated with surgery alone and 31% of the patients received neoadjuvant chemotherapy (CTx). Collection B was formed as the validation collection and comprised 320 patients selected using the same inclusion criteria as those used for Collection A. Among these, 28% of the patients received neoadjuvant CTx, and their detailed clinicopathological characteristics have been summarized in Tables 1 and 2.

The response to preoperative CTx was determined histopathologically and classified into three tumor-regression grades (TRGs) – TRG1b, TRG2, and TRG3 – which corresponded to <10%, 10–50%, and >50% residual tumor cells [18]. All patients were treated with chemotherapeutic regimes based on platinum/5-fluorouracil (5-FU) (Table 1). Further, all surgical approaches included an abdominal D2 lymphadenectomy [19].

The follow-up data were obtained from the tumor data management of the University Hospital of Augsburg, and the median follow-up was calculated using the inverse Kaplan–Meier method [20]. The primary endpoint of the study was overall survival (OS), which was defined as the time elapsed between the date of diagnosis and the death of the patient due to any cause. Patients with an OS of less than 30 days were excluded from the study.

The study was approved by the ethical committee of the Ludwig Maximilian University of Munich (reference: 20-0922) and was performed in accordance with the Declaration of Helsinki. According to the ethics vote of the institutional review boards of the Ludwig Maximilian University of Munich (reference: 20-0922), a declaration of consent was not necessary.

#### Definition and assessment of SARIFA

SARIFA was evaluated based on entire slide sections that covered about 220 mm<sup>2</sup>. We have defined SAR-IFA as an area located at the invasion front (IF) in which a tumor gland or a group of at least five tumor cells directly approach adipocytes without separating stroma. In addition, SARIFA was said to be present when tumor cells were directly adjacent to local tissue, such as the smooth muscle in muscularis propria, without a stromal reaction, such as fibroblastic proliferation, collagen formation, or a histiocytic reaction. We classified tumors that presented these characteristics as SARIFA-positive and the others as SARIFA-negative. Even if only a single SARIFA was present (e.g. a single tumor gland surrounded directly by inconspicuous adipose tissue), we classified the tumor as SARIFA-positive. The first (FA), third (TA), and last author (LA) independently assessed all tumors. They did not have access to each other's results or the clinical data. An exemplary tumor slide preselected by FA was used for assessment in each case. Further, any discrepant results were discussed using a multi-head microscope, and a consensus was reached. Example images are presented in Figure 1.

# Tissue microarray construction and TCGA classification

For the classification according to the TCGA, tissue microarrays (TMAs) were constructed. Tumors were

Table 1.	Clinicopathological	characteristics – Collection A.

Variable Median age (range), years Median follow-up (95% Cl), months		n = <b>160</b> 70.5 (30-95)		<b>SARIFA-positive (</b> <i>n</i> = <b>24)</b> 71 (40–87)		SARIFA-negative ( $n = 136$ ) 70.5 (30–95)		<i>P</i> value 0.100
		Sex						
	Female	58	36%	10	42%	48	35%	
<b>T</b>	Male	102	64%	14	58%	88	65%	0.004
i status	nT1/2	61	200%	1	40%	60	4.40%	<0.001
	pT1/2 nT3/4	90	50% 62%	23	4%0 960/o	60 76	44%0 560/o	
N status	p13/+	55	02-70	23	30%	70	30%	0.088
	Negative	58	36%	5	21%	53	39%	0.000
	Positive	102	64%	19	79%	83	61%	
Distant metastasis								0.009
	No	83	52%	7	29%	76	56%	
	Yes	67	42%	16	67%	51	38%	
	NA	10	6%	1	4%	9	7%	
Grading	Lever and a	54	2.40/	7	000/	47	250	0.682
	Low grade	54 104	34%	10	29%	4/	35%	
		104	05%0 10/6	10	67%0 40/o	00	10/2	
Vascular invasion		2	190	I I	4%0	I.	1.40	0 184
	Negative	145	91%	20	83%	125	92%	0.101
	Positive	15	9%	4	17%	11	8%	
Lymphovascular invasion	ı							0.047
	Negative	108	68%	12	50%	96	71%	
	Positive	52	33%	12	50%	40	29%	
Laurén classification								0.265
	Intestinal	90	56%	11	46%	79	58%	
1 P 0	Non-intestinal	70	44%	13	54%	57	42%	4 000
Localization	Drovimol	40	250%	C	250%	24	2506	1.000
	Non-provimal	40	25%0 75%	19	25%0 75%	34 102	25%	
R status	Non-proximal	120	7 3 %0	10	7 3%0	102	7 3 %0	0 389
it status	RO	141	88%	20	83%	121	89%	0.000
	R1	13	8%	3	13%	10	7%	
	Rx	6	4%	1	4%	5	4%	
TCGA								0.218
	EBV <sup>+</sup>	12	8%	0	0%	12	9%	
	MMRD	18	11%	2	8%	16	12%	
	GS	35	22%	4	17%	31	23%	
	CIN Nam alassifiable	46	29%	10	42%	36	26%	
	Non-classifiable	40	25%	/	29%	33	24%	
Death	INA	9	0%0	1	4%0	o	0%0	0.006
Death	No	81	51%	6	25%	75	55%	0.000
	Yes	79	49%	18	75%	61	45%	
nCTx								0.811
	No	110	69%	17	71%	93	68%	
	Yes	50	31%	7	29%	43	32%	
TRG								0.150
	1b	5	10%	0	0%	5	12%	
	2	19	38%	1	14%	18	42%	
CTx regime	3	26	52%	6	86%	20	47%	0.200
cix regime	Cis/Ox/5_EU or Con	12	260/2	0	00%	12	300%	0.206
	0x + 5-FU + Doc	19	38%	2	29%	17	40%	
	Cis + 5-FU + Epi	13	26%	4	57%	9	21%	
	Ox + Epi + Cap	4	8%	1	14%	3	7%	
	Others	1	2%	0	0%	1	2%	

P values are shown for difference between SARIFA-positive and SARIFA-negative tumors; values in bold are statistically significant.

5-FU, 5-fluorouracil; Cap, capecitabine; ClN, chromosomally instable; Cis, cisplatin; Doc, docetaxel; EBV<sup>+</sup>, EBV positive; Epi, epirubicin; GS, genomically stable; MMRD, mismatch repair deficient; nCTx, neoadjuvant CTx; Others, combination of Cis/Ox with other agents or crossover between different treatment regimens; Ox, oxaliplatin; Pac, paclitaxel; TCGA, The Cancer Genome Atlas; TRG, tumor regression grade.

classified in analogy to the TCGA classification [13] based on the immunohistochemical expression of PMS2, MSH6, E-cadherin, p53, and EBER-CISH [21,22]. The

TMA construction and classification according to the TCGA are described in more detail in the Supplementary materials and methods.

Variable Median age (range), years Median follow-up (95% Cl), months		n = 320	)	SARIFA-positive ( $n = 72$ )		SARIFA-negat	SARIFA-negative ( $n = 248$ )	
		69.0 (36	-91)	70.0 (38–91)		69.0 (36–90)		0.100
		62.0 (49	.6-74.4)	58.0 (26.5-8	9.5)	63.0 (49.0-77.	0)	0.912
Sex		(	,		,		-,	0.533
	Female	103	32%	21	29%	82	33%	
	Male	217	68%	51	71%	166	67%	
Tistatus	Marc	217	00 /0	51	7170	100	07 /0	~0.001
i status	nT1/2	02	200%	7	100%	95	340%	<0.001
	pT1/2	22	Z 3%0	CE /	00%	162	54%	
N status	p13/4	220	7190	05	30%0	105	00%0	0.004
IN Status	Negetius	117	270/-	10	2204	101	410/-	0.004
	Desitive	117	37%	16	22%	101	41%	
Distant materia	Positive	203	63%	56	78%	147	59%	-0.001
Distant metastasis		101	500/		0001	100	500/	<0.001
	No	161	50%	23	32%	138	56%	
	Yes	136	43%	45	63%	91	37%	
	NA	23	7%	4	6%	19	8%	
Grading								0.083
	Low grade	105	33%	17	24%	88	35%	
	High grade	210	66%	52	72%	158	64%	
	NA	5	2%	3	4%	19	8%	
Vascular invasion								0.017
	Negative	258	81%	51	71%	207	83%	
	Positive	62	19%	21	29%	41	17%	
Lymphovascular invasio	n							0.018
	Negative	181	57%	32	44%	149	60%	
	Positive	139	43%	40	56%	99	40%	
Laurén classification								0.648
	Intestinal	172	54%	37	51%	135	54%	
	Non-intestinal	148	46%	35	49%	113	46%	
Localization								0.087
	Proximal	90	28%	26	36%	64	26%	
	Non-proximal	230	72%	46	64%	184	74%	
R status		200	12.10	10	0170	101	7170	0.095
in Status	RO	265	830/	52	720/0	213	860/0	0.000
	R1	12	120/2	12	190/2	215	120%	
	Rv	12	10%	7	10%	23	20/2	
TCCA	INA	15	4%0	/	10%0	0	2%0	0.225
ICUA		10	40/-	4	<b>C</b> 0/ <del>-</del>	0	20/-	0.225
		12	4%	4 7	10%	0	3%	
		41	13%	/	10%	34	14%	
	GS CINI	/3	23%	12	17%	61	25%	
		95	30%	26	36%	69	28%	
	Non-classifiable	82	26%	20	28%	62	25%	
	NA	17	5%	3	4%	14	6%	
Death								0.004
	No	146	46%	22	31%	124	50%	
	Yes	174	54%	50	69%	124	50%	
nCTx								0.503
	No	230	72%	54	75%	176	71%	
	Yes	90	28%	18	25%	72	29%	
TRG								0.143
	1b	9	10%	0	0%	9	13%	
	2	22	24%	3	17%	19	26%	
	3	59	66%	15	83%	44	61%	
CTx regime								0.505
5	Cis/Ox/5-FU or Can	15	17%	2	11%	13	18%	
	0x + 5-FU + Doc	31	34%	6	33%	25	35%	
	Cis + 5-FU + Epi	31	34%	9	50%	22	31%	
	0x + Epi + Cap	11	12%	1	6%	10	14%	
	Others	2	2%	0	0%	2	3%	
				-				

Table 2. Clinicopathological characteristics - Collection B.

P values are shown for difference between SARIFA-positive and SARIFA-negative tumors; values in **bold** are statistically significant.

5-FU, 5-fluorouracil; Cap, capecitabine; CIN, chromosomally instable; Cis, cisplatin; Doc, docetaxel; EBV<sup>+</sup>, EBV positive; Epi, epirubicin; GS, genomically stable; MMRD, mismatch repair deficient; nCTx, neoadjuvant CTx; Others, combination of Cis/Ox with other agents or crossover between different treatment regimens; Ox, oxaliplatin; Pac, paclitaxel; TCGA, The Cancer Genome Atlas; TRG, tumor regression grade.

## NanoString GeoMx digital spatial profiling (DSP)

The DSP analysis is described in more detail in the Supplementary materials and methods. Using DSP, we performed a multiplexed and spatially resolved profiling analysis on exemplary SARIFA-positive and SARIFAnegative cases of TMAs (six SARIFA-positive and six



**Figure 1.** H&E images of SARIFA-positive and SARIFA-negative cases: SARIFA-positive cases (A, C, E: scale bar = 200  $\mu$ m; B, D, F: scale bar = 50  $\mu$ m) with tumor cells directly adjacent to local tissue, such as adjpocytes (A–D) or smooth muscle in muscularis propria (E, F), without a stromal reaction. SARIFA-negative case at the invasion front (G: scale bar = 200  $\mu$ m; H: scale bar = 50  $\mu$ m).

SARIFA-negative cases). DSP technology uses RNA detection probes with ultraviolet (UV) photocleavable indexing oligos for transcriptomic profiling within the selected regions of interest (ROIs). As shown in Figure 3A,B, based on fluorescence imaging to visualize the tumor cells and the corresponding H&E images, ROIs within the SARIFA-positive and SARIFA-negative areas were chosen for multiplex profiling. Sequencing libraries were generated as described previously [23] according to NanoString's GeoMx-NGS Readout Library Prep instructions (NanoString

Technologies, Inc, Seattle, WA, USA) and sequenced on an Illumina NovaSeq (Illumina Inc, San Diego, CA, USA).

Immunohistochemical protein expression of differentially expressed genes in DSP analysis and of proteins associated with fatty acid metabolism, hypoxia, and stem cell features

The immunohistochemical expression of FABP4, CD36, CD44, carbonic anhydrase IX, CD68, and Ki67

obtained from 30 SARIFA-positive and 30 SARIFAnegative cases was analyzed on 2 µm whole-slide sections using the primary antibodies listed in supplementary material, Table S1. Staining was performed on the BOND Rx platform using the BOND Polymer Refine Detection kit (Leica Biosystems, Nussloch, Germany). Adequate controls were used for quality control during staining (see supplementary material, Table S1). The staining was assessed separately for the IF and tumor center (TC). For CD36, CD44, and carbonic anhydrase IX, the staining was evaluated using the seven-tier semi-quantitative scoring system (immunoreactive score, IRS) proposed by Remmele and Stegner [24] while taking into account staining intensity and the percentage of positive tumor cells. For FABP4 and Ki67, the percentage of tumor cells positive was assessed. Finally, the abundance of macrophages was assessed through CD68 staining.

## Statistical analyses

Chi-squared tests were used for hypothesis testing of differences between relative frequencies. Continuous variables were compared using the Wilcoxon rank-sum test, while correlations were calculated using the Spearman test. Kaplan-Meier estimates of survival rates were compared using log-rank tests, and relative risks were estimated by hazard ratios (HRs) obtained from Cox proportional hazard models. Because overall patient survival was chosen as the study endpoint, an additional comparison of survival curves was performed using the Gehan-Breslow-Wilcoxon (GBW) test, as this gives more weight to early events that are more likely to be cancer-specific. Kappa statistics were used as a measure of inter-observer agreement [25]. Statistical analyses were performed using SPSS version 24 (IBM Corp., Armonk, NY, USA) and R version 4.0.3. Exploratory significance levels (two-tailed) of 5% were used for hypothesis testing. The DSP analysis was conducted using GeoMx Analysis Suite Version 2.2.0.111 (NanoString Technologies, Inc).

## Results

For Collection A, the mean age of the patients was 70.5 (range 30.0–95.0) years and the median follow-up time was 56.0 (range 44.8–67.2) months. In total, 49% of the patients died during the follow-up period. Detailed clinicopathological characteristics are summarized in Table 1. Overall, 24 (15%) cases were classified as SARIFA-positive, and 136 (85%) were deemed SARIFA-negative.

For Collection B, the mean age of the 320 patients at the time of diagnosis was 69.0 (range 36.0–91.0) years and the mean follow-up time was 62.0 (range 49.6–74.4) months. Of the 320 patients, 54% died during the follow-up period. Detailed clinicopathological characteristics are provided in Table 2. Overall, 72 (22.5%)

tumors were classified as SARIFA-positive, while 248 (77.5%) were classified as SARIFA-negative.

## SARIFA presents high inter-observer concordance

The first (FA), third (TA), and last author (LA) independently assessed all the tumors included in this study.

For Collection A, the overall kappa value obtained by all three observers was 0.738 (FA and TA: 0.727; FA and LA: 0.860; TA and LA: 0.636). For Collection B, the overall kappa value obtained by all three observers was 0.775 (FA and TA: 0.839; FA and LA: 0.763; TA and LA: 0.721).

## SARIFA and clinicopathological characteristics

The SARIFA-positive tumors were associated with several clinicopathological factors in both collections. An overview of this is presented in Tables 1 and 2 for Collection A and B, respectively. In both collections, SARIFA-positive tumors were associated with an advanced T-stage (p < 0.01 for each), distant metastases (p < 0.01 for each), and lymphovascular invasion (p < 0.05 for each). Additionally, in Collection B, the SARIFA-positive tumors were more frequently found to present positive lymph nodes (p < 0.01) and vascular invasion (p = 0.018). Further, no association with the molecular TCGA subtypes could be seen in Collection A or B (p = 0.218 and 0.225, respectively). As the interaction between adipose tissues and tumor cells has primarily been studied in the context of obesity research, we determined the relationship between SARIFA and patients' body mass index (BMI) in a subgroup of 60 cases (30 SARIFA-positive and -negative cases each). Here, SARIFA was associated with a lower BMI (median: 25 kg/m<sup>2</sup>) compared with non-SARIFA patients (median: 27 kg/m<sup>2</sup>) (p = 0.038).

## SARIFA and survival

For Collection A, patients who presented with SARIFApositive tumors had a significantly lower OS ( $p_{log}$  $_{rank} = 0.014$ , Figure 2A;  $p_{GBW} = 0.165$ ). SARIFA-positive patients had a median survival of 20.0 (11.0–29.0) months as opposed to 44.0 (17.7–70.3) months for SARIFA-negative cases. During the study period, 75% of the patients who had SARIFA-positive tumors died, whereas this was the case for 45% of the SARIFAnegative patients.

For Collection B, patients with SARIFA-positive tumors had a significantly lower OS ( $p_{log}$  rank +GBW < 0.001, Figure 2B). The SARIFA-positive cases showed a median survival of 15.0 (6.5–23.5) months, whereas patients with SARIFA-negative tumors had a median survival of 41.0 (21.8–60.2) months. During the study period, 69% of the patients who presented with SARIFA-positive tumors died, whereas this was the case for 50% of the SARIFA-negative patients.

Additionally, SARIFA positivity emerged as an independent negative prognostic factor (HR 1.638, 95% CI



Figure 2. Discrimination of patient survival based on SARIFA status. The Kaplan–Meier curves of patients with SARIFA-positive and SARIFAnegative tumors are shown. (A) Collection A (n = 160); (B) Collection B (n = 320). *P* value of log-rank test.

1.153–2.326, p = 0.006) for OS in Cox regression analyses, which were adjusted according to the known prognostic parameters (age, pT, pN, grading, and lymphovascular and vascular invasion). The results of these analyses are summarized in supplementary material, Table S2.

## Subgroup analysis

The subgroup analysis indicated the existence of a negative prognostic effect of SARIFA in primarily resected (HR 1.866, 95% CI 1.350–2.581, *p* < 0.001) and CTx patients (HR 2.003, 95% CI 1.145–3.506, p = 0.015). As shown in supplementary material, Table S3, SARIFA is a prognostic factor for intestinal or non-intestinal Laurén subtypes. The negative prognostic effect of SARIFA can especially be seen in non-proximal tumor localization. Regarding the pT stage, only advanced-stage (pT3/4) SARIFA-positive patients showed a worse OS (HR 1.618, 95% CI 1.200–2.180, p = 0.002), whereas no differences in terms of survival could be observed during the early stages (pT1/2) (HR 1.201, 95% CI 0.431-3.344, p = 0.726). Regarding the subgroups of TCGA, SARIFA was found to be a negative prognostic variable, especially in CIN (HR 1.756, 95% CI 1.106-2.788, p = 0.017) and non-classifiable tumors (HR 2.460, 95% CI 1.436–4.216, p = 0.001. Detailed information about this may be found in supplementary material, Table S3.

#### Differentially expressed genes identified using DSP

Using DSP technology, we identified genes that were significantly upregulated at the IF in SARIFA-positive cases compared with SARIFA-negative cases, as shown in Figure 3C. The most interesting upregulated genes in SARIFA-positive cases at the IF included *COL15A1* (p = 2.3569E-05), *FABP2* (p = 0.000226326), *FABP4* (p = 0.000763128), and *FGB* (p = 0.001792231). Figure 3 shows the differential expression of *COL15A1* (Figure 3D) and *FABP4* (Figure 3E) with the lowest

expression in the SARIFA-negative areas at the IF. *FAPB4* expression increases toward SARIFA-positive centers and again toward SARIFA-positive IFs. The expression of *COL15A1*, *FABP2*, *FABP4*, and *FGB* in the SARIFA-positive and SARIFA-negative areas is demonstrated in Figure 3F.

In pathway analyses, the most upregulated genes in the SARIFA-positive cases were found to be those associated with triglyceride catabolism, endogenous sterols, collagen chain trimerization, and fibrin clot formation.

## Immunohistochemical detection of the protein products of differentially expressed genes in DSP analysis, and of proteins associated with hypoxia and stem cell features

We wanted to confirm the differential expression of FABP4, which was identified using DSP analysis, at the protein level. FABP4 was almost exclusively expressed in the SARIFA areas, whereas it was totally absent in the TC, or in non-SARIFA cases at the IF or TC (IF SARIFA versus non-SARIFA p < 0.001; Figure 4A). Cancer cells acquire exogenous fatty acids through cell-surface acid translocase, which is also called CD36 [26,27]. CD36 was significantly upregulated at the IF (p < 0.001) in SARIFA cases and was almost absent or present at extremely low expression at the TC and in non-SARIFA cases (Figure 4B). To assess the number of tumor-associated macrophages (TAMs), we immunostained for CD68. We observed that the number of CD68<sup>+</sup> cells increased at the IF (p = 0.019) and in the TC (p = 0.060) in SARIFA cases compared with SARIFA-negative tumors (Figure 4F).

To verify whether hypoxia plays a role with regard to SARIFA, we assessed the expression of carbonic anhydrase IX. As shown in Figure 4G, no difference between SARIFA and non-SARIFA cases was observed. In Figure 4H, the expression of CD44 (a marker of stem cell features) is shown. Expression of CD44 was significantly lower at the IF in SARIFA cases (p < 0.001) and



**Figure 3.** Transcriptome analysis using DSP. (A, B) H&E slide of a TMA core of a SARIFA-positive tumor (A) and serial fluorescence image visualizing the tumor cells in the SARIFA-area. Based on fluorescence imaging, ROIs were chosen for multiplex profiling (B). (C) Volcano plot showing differentially expressed genes between SARIFA-positive and SARIFA-negative cases at the IF in DSP analysis. (D, E) *COL151A* (D) and *FABP4* (E) are significantly differentially expressed in SARIFA-negative, and SARIFA-positive TC and SARIFA-positive IF, showing the highest expression in tumor cells at the IF in SARIFA cases. (F) Heatmap demonstrating the expression (digital counts) of *COL15A1*, *FABP2*, *FABP4*, and *FGB* in all analyzed cases of non-SARIFA cases at the IF, and SARIFA cases in the TC and at the IF.

TC (p = 0.001). Based on the proliferation-associated marker Ki67, no difference was seen between the SAR-IFA and non-SARIFA cases with regard to proliferation activity (Figure 4I).

The expression of FABP4 at the IF showed a significant inverse correlation with CD44 at the IF ( $\rho = -0.446$ , p < 0.001) and in the TC ( $\rho = -0.476$ , p < 0.001). FABP4 IF was found to be positively



**Figure 4.** Immunohistochemical protein expression of differentially expressed genes in DSP analysis and of proteins associated with hypoxia and stem cell features. (A–C, F–I) Boxplots showing expression of FABP4 (A) and CD36 (B); BMI (C); (F) number of CD68<sup>+</sup> cells; expression of carbonic anhydrase IX (G), CD44 (H), and Ki67 (I); and examples of immunohistochemical staining. (D, E) Correlation matrix of the proteins at the IF and the TC.

correlated with the amount of CD68<sup>+</sup> cells at the IF ( $\rho = 0.316$ , p = 0.019) or in the TC ( $\rho = 0.388$ , p = 0.003) and with CD36 expression at the IF ( $\rho = 0.535$ , p < 0.001) (Figure 4D,E).

## Discussion

So far, only a few prognostic and therapeutic biomarkers have been identified for gastric cancer. To date, the most important therapeutic marker in gastric carcinoma is HER2 overexpression [28]. In addition, MSI status and high PD-L1 expression are independent positive prognostic factors in gastric carcinoma [14,29,30], while aberrant E-cadherin expression is considered an unfavorable prognostic factor, and even a negative predictive factor for chemotherapy response [31]. Additionally, MET amplification and overexpression are thought to play a crucial role in gastric carcinogenesis, but this is yet to be found as a predictive factor for the response to anti-MET antibodies [14]. In this study, we present SARIFA as a new histological prognostic biomarker in gastric cancer that is easy to assess during routine diagnostic assessment. When evaluating SARIFA, there is no need for additional immunohistochemical staining or complicated evaluation schemes, such as those used for the evaluation of protein-based TCGA classification [21,22], tumor budding, the TSR [10–12], or the aforementioned prognostic biomarkers.

In addition, the criteria for SARIFA positivity are defined simply, which is reflected in the considerably low inter-observer variability with kappa values of 0.74 (Collection A) and 0.78 (Collection B) across all three observers. The presence of a single tumor cell next to inconspicuous adipocytes without the surrounding stroma is sufficient for the identification of SARIFA positivity.

SARIFA positivity was found to be a significant prognostic factor in both collections and emerged as an independent negative prognostic factor for overall survival in Collection B. As the majority of SARIFA cases are in advanced T-stage (90%), SARIFA by its nature is mainly a prognostic marker in already advanced cases. In summary, SARIFA combines a high degree of reproducibility, low additional effort, and high clinical relevance. Recently, Wulczyn *et al* [32] identified a morphologically similar phenomenon in colon carcinoma using a deep learning system.

To provide hints with regard to the underlying tumorpromoting mechanisms associated with SARIFA positivity, we conducted DSP. Among the most upregulated pathways at the IF in SARIFA-positive cases, compared with the negative cases, were those of collagen chain trimerization and fibrin clot formation. *COL15A1* was highly differentially expressed when comparing SARIFA-positive and SARIFA-negative cases. In the positive cases, its expression was also higher at the periphery than in the tumor's center. At first glance, this seems contradictory, as SARIFA-positive cases do not induce a stromal reaction and are characterized by the absence of fibrous stroma between adipocytes and tumor. COL15A1 encodes the alpha chain of type XV collagen, which represents one component of the basement membrane (BM) and is thought to function as an adhering component between the BM and the surrounding matrix [33]. Karppinen et al [34] detected collagen XV in the tumor stroma during late stages of tumor progression in squamous cell carcinoma. In our DSP analysis, only the expression of tumor cells was assessed and not explicitly of the stroma. Thus, the high expression in the stroma is not an explanation for COL15A1 expression in the tumor cells of SARIFA-positive cases. One could speculate that COL15A1 is highly expressed because of the synthesis of a non-functional protein, resulting in inadequate collagen synthesis. But this is highly speculative and needs to be investigated in further studies. SARIFA-positive cases were associated with an increased number of CD68-positive macrophages. TAMs can polarize toward M2 macrophages that enhance angiogenesis, tumor invasion, and immune suppression [35,36]. It is therefore probable that macrophages play a role in preventing the tumor glands from being entrapped by desmoplastic fibers and induce a reduced response of the immune system in SARIFA tumors. However, the characteristics of TAMs and the polarization between M1 and M2 macrophages that are known so far are based on in vitro experiments and cannot be transferred without limitations *in vivo*. The exact mechanisms that lead to this phenomenon and the role of TAMs, which seem to have a great heterogeneity and comprehensive functions, remain to be elucidated [37].

Interestingly, other pathways that were significantly upregulated in SARIFA-positive cases at the IF were those of triglyceride catabolism and endogenous sterols. Certain studies in the context of obesity research have investigated the mechanisms of tumor-associated adipocytes using cell culture experiments [16,26,38]. Wen et al [39] report that colon cancer cells uptake fatty acids secreted by adipocytes and that adipocytes activate autophagy to support cancer cell survival by altering homeostasis and cellular metabolism in cancer cells. Among the most differentially expressed genes were fatty acid-binding proteins 2 and 4 (FABP2 and FABP4). Fatty acid-binding proteins (FABPs) facilitate fatty acid transport to different cell organelles [40]. This seems to be an interesting starting point for further investigations to explain the underlying concepts of SARIFA and its association with adverse prognosis and characteristics.

To confirm the differential expression of FABP4 at the protein level, we used immunohistochemistry. We found that FABP4 was expressed almost exclusively in the tumor cells of SARIFA areas, whereas no expression was found in the TC or at the IF of SARIFA-positive and SARIFA-negative cases. Our finding regarding FABP4 expression in SARIFA areas supports the hypothesis that tumor cells gain energy through tumor fatty acid oxidation [41]. In addition, higher expression of CD36, which enables tumor cells to take up free fatty acids [26,27], was observed in the SARIFA areas. It was also observed that CD36-positive tumor cells represent a metastasisinitiating cell population [42,43]. Thus, fatty acid oxidation provides an efficient energy source for the tumor [44]. The inhibition of this pathway could be an attractive therapeutic option. In this respect, the transport of acyl-CoAs into mitochondria by carnitine palmitoyl transferase 1 (CPT1) represents a potential target [45].

The interaction between adipocytes and tumor cells has been studied mainly in obesity research to further address the questions of why obese persons have an increased incidence of cancer with a poorer prognosis [16,38]. However, we did not observe an association between SARIFA positivity and higher BMI; on the contrary, SARIFA was even associated with lower BMI in patients (Figure 4C). Thus, obesity does not appear to be the trigger for tumor progression here. Interestingly, the tumor cells also did not appear to be affected by hypoxia, as indicated by the low expression of the hypoxia marker carbonic anhydrase IX. Further, stem cell features also do not seem to play a role in tumor progression, and there even seems to be a lower expression of the stem cell marker CD44 in SARIFA cases [46–49]. Rather, the tumor cells appear to be able to proliferate unimpeded and invade the tissue. In doing so, they benefit from their contact with adipocytes. It is possible that tumor-associated macrophages induce a reduced host tissue response. Thus, the term 'stroma areactive' describes the observed biological process quite well. The indications of SARIFA being the morphological correlate of a certain tumor biology implicate their possible role as a predictive marker for a therapy targeting TAMs or overcoming the therapy resistance created by tumor-associated fat [16,35]. In this regard, our study has certain limitations, which are mainly related to its retrospective nature. Our study has to be considered an exploratory analysis, and the results have to be validated using independent prospective cohorts.

This study presents SARIFA (Stroma Areactive Invasion Front Areas) as an extremely promising histomorphology-based prognostic factor that is related to tumor-promoting adipocytes in gastric cancer. Further studies will verify whether the results of this retrospective, single-center study can be confirmed and whether SARIFA can serve as a universal prognostic indicator for solid cancers.

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#### Author contributions statement

BM, BG and BMa contributed to the study's conception and design. MG, DV, AV, DK, SD, AP, GS and BG

contributed to the data acquisition process. CD, SS, BM and BG contributed to the analysis and interpretation of data. All authors revised the article critically, contributed to it with reflective improvements, and approved the final version.

#### Data availability statement

The datasets generated during DSP analysis are available at https://doi.org/10.6084/m9.figshare.16615951.

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References 50–52 are cited only in the supplementary material.

#### SUPPLEMENTARY MATERIAL ONLINE

Supplementary materials and methods

 Table S1. Antibodies and dilutions used

 Table S2. Cox regression analyses

Table S3. Subgroup analyses: Cox regression analyses