Basal microvilli define the metabolic capacity and lethal phenotype of pancreatic cancer

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Abstract

Apical microvilli of polarized epithelial cells govern the absorption of metabolites and the transport of fluid in tissues. Previously, we reported that tall and dense basal microvilli present on the endothelial cells of pancreatic cancers, a lethal malignancy with a high metabolism and unusual hypomicrovasularity, contain nutrient trafficking vesicles and glucose; their length and density were related to the glucose uptake of pancreatic cancers in a small-scale analysis. However, the implications of basal microvilli on pancreatic cancers are unknown. Here, we evaluated the clinical implications of basal microvilli in 106 pancreatic cancers. We found that basal microvilli are a dominant change in pancreatic cancers. The presence of longer and denser basal microvilli on the microvessels in pancreatic cancer tissues positively correlated with increased glucose uptake and higher metastatic (or invasive) and proliferative potentials of neoplastic cells and vice versa. Clinically, postoperative patients with longer and denser basal microvilli were more prone to unfavorable pathological characteristics and dismal prognoses. They were even more refractory to adjuvant therapy than those with shorter and thinner basal microvilli were. Our findings show that basal microvilli define the metabolic capacity and lethal phenotype of pancreatic cancers.

Keywords: basal microvilli; pancreatic cancer; glucose uptake; lethal phenotype; microvessel

Introduction

Apical microvilli of the polarized epithelia govern the absorptive and transporting capacity of metabolites and fluid in human tissues. Abundant apical microvilli are present in the epithelial cells of the small intestine and renal proximal tubule, where massive exchanges of metabolites and fluid occur [1–4]. Pancreatic cancer is a highly metabolic and almost incurable tumor with unusual hypomicrovasularity and dense stroma [5–7]. Dense desmoplastic stroma and hypomicrovasularity in the pancreatic cancer milieu block the delivery of therapeutic agents to the tumor and lead to a grim outcome for the patient [8,9]. However, nutrients, including glucose, glutamine, and albumin, quickly reach the tumor milieu and efficiently support tumor growth and metabolism [8,10]. Previously, we described a recently recognized anatomical structure, basal microvilli, which are preferentially present on the basal surface of polarized endothelial cells in pancreatic cancer tissues; they traverse dense stroma and are close to neoplastic cells [11]. Basal microvilli contain pinocytic and macropinocytic vesicles and glucose and do not depend on angiogenesis [11,12]. Principally, densely packed and long microvilli infer the high exchanging capacity of epithelial tissue [1,13]. Thus, the dynamics of basal microvilli might define the glucose metabolic capacity and lethal phenotype of pancreatic cancers.

Apical microvilli increase the surface area and enrich membrane-associated molecules for absorbing nutrients [2]. Apical microvilli on epithelial tissues are highly efficient in selectively exchanging nutrients and waste. Atrophy of the intestinal microvilli causes life-threatening watery diarrhea in microvillus inclusion disease [14,15]. Microvilli of syncytiotrophoblast cells in
the placenta also facilitate bidirectional feto-maternal exchanges [13]. We found that macrophagic, pino-
cytic, Glut1-positive vesicles, and glucose appear in
the lumen of basal microvilli [11,12]. Therefore, basal
microvilli might compensate for the hypomicrovascular-
ity in pancreatic cancer tissues and selectively deliver nutri-
ents such as glucose to the tumor milieu rather than drugs
such as nucleoside analogs. By analyzing seven patient
samples, we found that longer and denser basal microvilli
are related to higher glucose uptake in pancreatic cancer
[11]. Higher glucose uptake in pancreatic cancer indicates
a poor prognosis after surgery [16–19]. Therefore, the
dynamics of basal microvilli might reflect the strengthening
of glucose metabolic capacity, metastasis, aggressiveness,
and chemoresistance of neoplastic cells in pancreatic can-
cers and predict the patient’s outcome.

This study analyzed the basal microvilli parameters in
106 consecutive pancreatic cancer patients and the
implications of basal microvilli in disease progression
after surgery. We aimed to determine the implications
of basal microvilli on the biology of neoplastic cells,
including nutrient uptake, proliferation, and metastatic
(or invasive) potential, and to define the relationships
between the basal microvilli parameters and the clinico-
pathological characteristics of pancreatic cancer patients
and the clinical values of basal microvilli for predicting
postoperative chemoresistance and prognosis.

**Patients and methods**

**Patient selection, data collection, and ethics**

This prospective study enrolled 106 consecutive pancreatic
cancer patients who underwent radical surgical resection
in two medical centers from May 2013 to December 2018.
Patients were excluded if neoadjuvant chemotherapy had
been administered preoperatively. The ethics committee
of Huashan/Zhongshan Hospital of Fudan University
approved the study and informed consent was obtained
from each patient. We have included the seven patients in
our previous publication [11]. The basal microvilli parame-
ters in tumor tissues were measured postoperatively, along-
side postoperative pathological assessments. Standard
demographic and clinicopathological data were obtained
from the Pancreatic Tumor Registry and analyzed simulta-
neously. The basal microvilli parameters included their
length and density, which were used to test the correlations
of basal microvilli with clinicopathological features and
prognosis. After the operation, patients were referred to
the department of oncology for adjuvant therapy.

**Immunofluorescence staining and antibodies**

Details of the antibodies and software used are provided
in supplementary material, Table S1.

Tissue collection and staining were conducted accord-
ing to a method described previously [11]. In brief, thin
slices of tumor samples were obtained and put into fresh
4% paraformaldehyde (PFA) for fixation. The tissue was
dehydrated in 30% sucrose, frozen, and cryosectioned
(45 μm). Sections were incubated in a blocking buffer
and then incubated in a primary antibody solution over-
night at 4 °C. Next, the sample was incubated for 1 h in
the appropriate secondary antibody solution containing
DAPI [donkey anti-rabbit Alexa 488 (Cat # A32790,
1/1000 dilution; Thermo Fisher, Rockford, IL, USA) or
donkey anti-mouse Alexa 555 or Cy3 (Cat # A-31570,
1/1000 dilution; Thermo Fisher; or Cat # 715-167-003,
1/1000 dilution; Jackson Lab, Philadelphia, PA,
USA)]. Finally, the immunostained sections were
mounted on slides. Zeiss 710 or 880 confocal micros-
copy (ZEISS, Jena, Germany) with Z-stack and tile
scanned images was performed. CD34 antibody (Cat #
ab8536, 1/1000 dilution; Abcam, Cambridge, MA,
USA) and cytokeratin-19 (CK-19) antibody (Cat #
ab52625, 1/500 dilution; Abcam) or GLUT-1 antibody
(Cat # ab115730, 1/200 dilution; Abcam) were used to
stain the microvessels and neoplastic cells, respectively.

**PET-CT scanning**

The scanning settings for the VCT 64 PET/CT scanner
(GE Medical Systems, Waukesha, WI, USA) were
140 kV and 200 mA. Each patient was injected with
0.1 mCi/kg body weight of 18F-FDG as a tracer. Three-
dimensional whole-body PET acquisitions were per-
formed at three time points with nominal start times at
40, 65, and 90 min after the injection. The filtered back-
projection (FBP) method was applied for image recon-
struction. The maximum standardized uptake value
(SUVmax) was quantified on the PET/CT images by
two experienced nuclear radiologists.

**Measurements of the basal microvilli parameters**

ImageJ software (Fiji, NIH, Bethesda, MD, USA) was used
to measure and analyze basal microvilli parameters as pre-
viously described [8]. We quantified the basal microvilli’s
length and density on the microvessels of pancreatic cancer
in a 4- to 15-image Z-stack taken using a 63× objective.
The mean/median values of length and density were con-
sidered as the basal microvilli parameters of patients
(sup-
plementary material, Table S2). The basal microvilli
index was calculated as \( \pi \times 1 \times \text{length} \times \text{density} \)/100.
Imaris 9.5 (Bitplane AG, Zürich, Switzerland) was used
to visualize the surface.

**Patient follow-up**

MRI/CT or PET-CT scanning, serum CA 19-9, clinical
symptoms, and physical examinations were documented
for postoperative surveillance. Follow-up information
was collected from the Pancreatic Tumor Registry, Fudan
University. Radiological examinations were performed
every 6 months or any time the patients had symptoms
related to signs of recurrence. Recurrence was defined as
radiological evidence of intra-abdominal soft tissue around
the surgical site or distant metastasis with or without CA
19-9 > 37 U/ml during postoperative surveillance. The
duration of overall survival (OS) was calculated from
the date of surgery to the date of tumor-specific death or the last follow-up visit date. Relapse-free survival (RFS) was calculated as the time from surgery to the time of recurrence.

Chemotherapy and chemoradiotherapy

Patients received gemcitabine (1000 mg/m² over 30 min) on days 1, 8, and 15 of a 28-day cycle, or gemcitabine (1000 mg/m² over 30 min) on days 1 and 8 with concurrent S-1 (40–60 mg/m²) orally twice daily on days 1–14 of a 21-day cycle. The adjuvant chemoradiotherapy dose was 50.4 Gy in 25–28 fractions and was given concurrently with S-1 (40 mg) orally twice daily during the radiation day.

Statistical analyses

The statistical analyses were performed using SPSS version 21.0 (SPSS Inc, Chicago, IL, USA) and R version 3.3.0 (http://www.r-project.org/). Pearson or Spearman correlation analysis was also performed to compare the correlations. Receiver operating characteristic (ROC) curves were used to evaluate the prognosis prediction of basal microvilli parameters. Survival curves were estimated according to the Kaplan–Meier method and were compared using the log-rank test. A Cox proportional hazards model was utilized for univariate analysis.

Results

General data

One hundred and six consecutive patients from two pancreatic cancer centers were recruited in this study (Table 1). Sixty-two samples (58.5%) were collected from the Department of Pancreatic Surgery, Zhongshan Hospital between May 2013 and December 2018; 44 (41.5%) were collected from the Pancreatic Cancer Center of Huashan Hospital between December 2016 and December 2018. The longest duration of follow-up was 1187 days, and the shortest was 133 days. The median follow-up time was 419 days. The median RFS time was 298/C6 28 days (1/C6 2 SEM), and the median OS was 585/C6 43 days (1/C6 2 SEM). Ninety-one patients (85.9%) relapsed after radical surgery, and 57 patients (53.8%) died during the follow-up period (Table 1). Resection margins, including transection and

| Table 1. Clinicopathological characteristics of the pancreatic cancer patients. |
|------------------|------------------|------------------|------------------|
| Clinical variables | Count (%) | RFS univariate | OS univariate |
| Gender Female | 44 (41.5) | 0.54 | 1.140 (0.748–1.738) | 0.785 | 0.928 (0.541–1.592) |
| Male | 62 (58.5) | 0.33 | 0.799 (0.507–1.259) | 0.718 | 0.899 (0.502–1.607) |
| Age <60 30 (28.3) | 0.26 | 1.286 (0.828–1.998) | 0.028 | 1.857 (1.069–3.228) |
| ≥60 76 (71.7) | 0.00 | 2.217 (1.408–3.490) | 0.008 | 2.261 (1.235–4.141) |
| Tumor stage T1 + T2 67 (66.8) | 0.14 | 1.512 (0.878–2.606) | 0.007 | 1.420 (1.103–1.830) |
| T3 + T4 39 (38.3) | 0.81 | 1.051 (0.668–1.672) | 0.48 | 0.821 (0.475–1.419) |
| RLM Yes 74 (69.8) | 0.51 | 1.158 (0.746–1.798) | 0.013 | 1.964 (1.150–3.553) |
| No 32 (30.2) | 0.00 | 2.084 (1.279–3.396) | 0.002 | 2.720 (1.463–5.056) |
| TNM I + II 89 (84.0) | 0.03 | 1.824 (1.057–3.146) | 0.133 | 1.660 (0.857–3.217) |
| III + IV 17 (16.0) | 0.91 | 1.036 (0.561–1.915) | 0.715 | 1.151 (0.540–2.452) |
| PNI Yes 74 (69.8) | 0.92 | 1.023 (0.648–1.615) | 0.818 | 1.072 (0.592–1.940) |
| No 32 (30.2) | 0.55 | 1.134 (0.748–1.719) | 0.194 | 1.430 (0.837–2.446) |
| MVI Yes 37 (34.9) | 0.01 | 3.312 (1.971–5.565) | <0.001 | 3.565 (1.809–7.024) |
| No 69 (65.1) | 0.00 | 3.312 (1.971–5.565) | <0.001 | 3.565 (1.809–7.024) |
| Tumor vascular thrombus Yes 27 (25.5) | 0.00 | 2.084 (1.279–3.396) | 0.002 | 2.720 (1.463–5.056) |
| No 79 (74.5) | 0.00 | 2.084 (1.279–3.396) | 0.002 | 2.720 (1.463–5.056) |
| Resectional margin R1 17 (16.0) | 0.91 | 1.036 (0.561–1.915) | 0.715 | 1.151 (0.540–2.452) |
| R0 89 (84.0) | 0.92 | 1.023 (0.648–1.615) | 0.818 | 1.072 (0.592–1.940) |
| CA 19-9 Positive 92 (86.8) | 0.55 | 1.134 (0.748–1.719) | 0.194 | 1.430 (0.837–2.446) |
| Negative 14 (13.2) | 0.00 | 3.312 (1.971–5.565) | <0.001 | 3.565 (1.809–7.024) |
| CEA Positive 31 (29.2) | 0.34 | 1.134 (0.748–1.719) | 0.194 | 1.430 (0.837–2.446) |
| Negative 75 (70.8) | 0.00 | 3.312 (1.971–5.565) | <0.001 | 3.565 (1.809–7.024) |
| Ki67 index <30% 46 (43.8) | 0.55 | 1.134 (0.748–1.719) | 0.194 | 1.430 (0.837–2.446) |
| ≥30% 59 (56.2) | 0.00 | 3.312 (1.971–5.565) | <0.001 | 3.565 (1.809–7.024) |
| PET-CT <6.5 34 (32.1) | 0.00 | 2.084 (1.279–3.396) | <0.001 | 3.565 (1.809–7.024) |
| >6.5 40 (37.7) | 0.00 | 2.084 (1.279–3.396) | <0.001 | 3.565 (1.809–7.024) |
| NA 32 (30.1) | 0.33 | 0.675 (0.308–1.476) | 0.45 | 1.427 (0.567–3.593) |
| Adjuvant therapy Chemo 87 (82.1) | 0.91 | 1.036 (0.561–1.915) | 0.715 | 1.151 (0.540–2.452) |
| Chemo + radio 10 (9.4) | 0.92 | 1.023 (0.648–1.615) | 0.818 | 1.072 (0.592–1.940) |
| NA 9 (8.5) | 0.55 | 1.134 (0.748–1.719) | 0.194 | 1.430 (0.837–2.446) |
| Relapse Yes 91 (85.8) | Median DFS: 298 ± 28 |
| No 15 (14.2) | 0.00 | 3.312 (1.971–5.565) | <0.001 | 3.565 (1.809–7.024) |
| Survival Alive 49 (46.2) | Median OS: 585 ± 43 |
| Died 57 (53.8) | 0.00 | 3.312 (1.971–5.565) | <0.001 | 3.565 (1.809–7.024) |

MVI, major vascular invasion; PNI, perineural invasion; RLM, regional lymph metastasis. Bold font indicates statistically significant values.
did not receive any adjuvant treatment (Table 1). Out concurrent S-1 or radiotherapy, and nine patients underwent surgery, including gemcitabine with or without concurrent S-1 or radiotherapy, and nine patients (91.5%) received chemotherapy-based treatment after surgery, including gemcitabine with or without concurrent S-1 or radiotherapy, and nine patients (8.5%) did not receive any adjuvant treatment (Table 1).

Basal microvilli are distinct characteristics of the microvasculature in pancreatic cancer

Basal microvilli are tiny, hair-like endothelial projections on the surface of the basal side of endothelia and can be visualized by high-resolution 3D confocal scans [11]. We have reported the patterns of basal microvilli on the microvessel segments in pancreatic cancers [11]. The depth and width of basal microvilli on the entire microvasculature nets in pancreatic cancer have not been reported. To see the basal microvilli patterns on the entire microvasculature nets of pancreatic cancer tissues, we visualized the whole networks of basal microvilli microvessels throughout the tumor region’s non-cancerous tissue by an extensive area stitching technique after CD34 and CK-19 antibody immunostaining in 45-μm-thick sections. We observed that the basal microvilli were distinct features and systemic changes in the pancreatic cancer microvasculature system (Figure 1A), and were mostly enriched in the smaller branches of the microvasculature networks (Figure 1B). Consistent with our previous studies [8], basal microvilli also existed on the microvasculature of the invasive frontier of the tumor (Figure 1A). Surprisingly, the microvessel networks in pancreatic islets that were surrounded by neoplastic cells did not present basal microvilli, but the surrounding microvessels in the tumor region presented with long and dense basal microvilli (supplementary material, Figure S1A). We also observed that extremely long, thick, and dense basal microvilli presented on the primary pancreatic cancer tissue microvasculature with distant metastasis to the liver (supplementary material, Figure S1B). These findings further indicated that basal microvilli were tumor-specific and indicated the metabolic capacity of tumors rather than healthy pancreatic tissues, reflecting the metastatic and invasive potential of neoplastic cells in pancreatic cancers.

Longer and denser basal microvilli are related to the increased glucose uptake, higher metastatic or invasive and proliferative potential of neoplastic cells in pancreatic cancer

To analyze these basal microvilli on the microvasculature in pancreatic cancers, we used immunofluorescence for CD34 and glucose transporter-1 (GLUT-1) or cytokeratin-19 (CK-19) on five to ten cryosections cut at 45 μm thickness. After staining, we scanned the slices using a 63x objective to find microvessels with basal microvilli, and compared the length and density of the basal microvilli on different microvessels. After finding the microvessels with longer and denser basal microvilli compared with other microvessels, we scanned 4–15 images from each patient’s tumor tissue, used 3D reconstruction, and then measured the length and density of the basal microvilli. Among 106 patients, imaging showed that the microvasculature presented basal microvilli in 103 patients, and that the microvasculature in three patients did not present basal microvilli. We previously reported that an abundance of basal microvilli was closely correlated with increased PET-SUVmax values in seven patients [11]. In this cohort, 74 patients had PET-CT scanning before surgery. Our correlation analysis showed that longer and denser basal microvilli significantly correlated with a higher PET-SUVmax value (Figure 2A). The higher the PET-CT SUVmax value of the tumor tissue, the longer and denser were the basal microvilli that presented in the microvasculature of the same region (Figure 2A). Moreover, we observed that a ‘flourishing’ basal microvilli pattern with widespread branches from the microvasculature in pancreatic cancer tissues reflected extremely high PET-CT SUVmax.
Figure 2. Longer and denser basal microvilli in pancreatic cancer indicate an elevated glucose uptake. (A) Typical patterns of basal microvilli on the microvasculature segments of nine pancreatic cancers with differential PET-CT SUVmax values. A CD34 antibody was used to stain microvessels. All images were taken from a Z-stack and are shown as maximum intensity projection. Scale bar: 20 μm. (B) The relationships of the mean/median values of basal microvilli length, density, and index with tumor PET-CT SUVmax values in pancreatic cancer patients. Spearman’s correlations were used to estimate the correlation efficiency and statistical significance.

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we used the microvilli was assessed by density and length; therefore, $r = 0.737$ (median). The cellular surface of the basal microvilli were more prone to having regional lymph node metastasis, advanced TNM stage, tumor thrombus, and a high Ki67 index (Table 2). The basal microvilli length was also associated with margin-positive resection (R1) and major vascular invasion (Table 2). Univariate analysis showed that regional lymph node metastasis, advanced TNM stage, tumor thrombus, margin-positive resection, and major vascular invasion were significant risk factors in this cohort (Table 1). These results demonstrated that longer and denser basal microvilli in pancreatic cancer indicated higher metastatic or invasive and proliferative potential in neoplastic cells.

Longer and denser basal microvilli in pancreatic cancer are related to the dismal prognosis

Pieces of evidence suggest that pancreatic cancer patients with high PET-CT SUVmax values and lymph metastasis have a dismal prognosis [17,18,20]. In our previous work, we did not explore the clinical implications of basal microvilli. After obtaining the patients’ follow-up data, we noticed that pancreatic cancer patients with a shorter relapse-free survival (RFS) have longer and denser basal microvilli on the tumor microvasculature, and vice versa (Figure 3A). Thus, basal microvilli may have the ability to predict relapse or death of pancreatic cancer patients. To test if basal microvilli parameters can predict relapse or death of pancreatic cancer patients, we carried out an ROC curve analysis. The results showed that all basal microvilli parameters, including density, length, and index, could predict the relapse or death of pancreatic cancer patients, and all the AUC (area under curves) values were higher than 0.649 (Figure 3B, C). We further analyzed the hazard ratios and survival functions of basal microvilli parameters by Cox regression and Kaplan–Meier methods. The analysis showed that patients with longer and denser basal microvilli had both significantly lower RFS and significantly lower OS days than patients with shorter and sparser basal microvilli [Figure 4A; supplementary material, Figure S3A, B; length: shorter versus longer, 419 versus 202 days (RFS); 998 versus 410 days (OS); density: thinner versus denser, 400 versus 233 days (RFS); 751 versus 467 days (OS)]. The hazard ratios of longer basal microvilli for RFS and OS rates were extremely high [supplementary material, Figure S3A, B; supplementary material, Figure S3A, B; length: 751 versus 467 days (OS)].

Table 2. Correlations of the basal microvilli parameters with clinicopathological variables in the pancreatic cancer patients.

|                       | BMD1       | BMD2       | BML1       | BML2       | BMI1       | BMI2       |
|-----------------------|------------|------------|------------|------------|------------|------------|
| Gender, male versus female | $r = 0.1$ | $p = 0.3$  | $r = 0.1$  | $p = 0.3$  | $r = 0.1$  | $p = 0.3$  |
| Age, >60 versus <60   | $r = -0.1$ | $p = 0.5$  | $r = -0.1$ | $p = 0.5$  | $r = -0.1$ | $p = 0.5$  |
| Tumor size            | $r = -0.3$ | $p = 0.1$  | $r = -0.3$ | $p = 0.1$  | $r = -0.3$ | $p = 0.1$  |
| RLM (yes versus no)   | $r = 0.1$  | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  |
| TNM stage             | $r = 0.9$  | $p < 0.001$| $r = 0.9$  | $p < 0.001$| $r = 0.9$  | $p < 0.001$|
| PNI (yes versus no)   | $r = 0.5$  | $p = 0.1$  | $r = 0.5$  | $p = 0.1$  | $r = 0.5$  | $p = 0.1$  |
| MVI (yes versus no)   | $r = 0.2$  | $p = 0.5$  | $r = 0.2$  | $p = 0.5$  | $r = 0.2$  | $p = 0.5$  |
| Serum albumin levels  | $r = 0.3$  | $p = 0.1$  | $r = 0.3$  | $p = 0.1$  | $r = 0.3$  | $p = 0.1$  |
| Fasting blood glucose | $r = 0.2$  | $p = 0.5$  | $r = 0.2$  | $p = 0.5$  | $r = 0.2$  | $p = 0.5$  |
| CA 19-9 (positive versus negative) | $r = 0.1$ | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  |
| CEA (positive versus negative) | $r = 0.1$ | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  |
| Ki67 index            | $r = 0.1$  | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  |
| Resection margin (R1 versus R0) | $r = 0.1$ | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  | $r = 0.1$  | $p = 0.5$  |

Spearman correlation, $r$, correlation efficiency; fasting blood glucose, before surgery; serum albumin levels, before surgery; tumor size, the largest diameters. BMD, basal microvilli density; BML, basal microvilli length; BMI, basal microvilli index; 1, mean; 2, median. Bold font indicates statistically significant values.
The hazard ratio of longer basal microvilli for OS and RFS rates was higher than that of regional lymph node metastasis, histopathological grade, TNM stage, peripheral neuronal invasion, major vascular invasion, Ki67 index, serum CA 19-9 and CEA values, and was almost equal to that of the SUVmax value on PET (Table 1). These findings showed that basal microvilli parameters, such as length and density, indicate the prognosis of pancreatic cancer patients after surgery.

Longer and denser basal microvilli in pancreatic cancer are refractory to adjuvant therapy

In our cohort, 94 patients with basal microvilli received adjuvant therapy, including chemotherapy or...
chemoradiotherapy, whereas nine patients did not receive any adjuvant treatment. In the adjuvant therapy subgroup, patients with longer and denser basal microvilli were more prone to have dismal outcomes than patients with shorter and sparser basal microvilli (Figure 5A,B and supplementary material, Figure S4A,B). In the adjuvant therapy subgroup, survival analysis also verified that patients with longer and denser basal microvilli had both significantly lower OS and significantly lower RFS rates than those with shorter and sparser basal microvilli [length: longer versus shorter, 217 versus 390 days (RFS) and 412 versus 998 days (OS); density: denser versus thinner, 237 versus 400 (RFS) and 467 versus 751 (OS)]. We observed that the median RFS and OS of pancreatic cancer patients with longer basal and denser microvilli were shorter than those of patients with shorter and thinner basal microvilli in the no-treatment subgroup [supplementary material, Figure S5; length: longer versus shorter, 116 versus 434 days (RFS) and 146 versus 585 days (OS); density: denser versus thinner, 112 versus 400 (RFS) and 112 versus 565 (OS)]. However, this analysis was carried out in a cohort of just nine patients (3+6) and the data are not significant. This finding suggested that basal microvilli parameters also indicate cancer cells’ sensitivity to adjuvant therapies, including chemotherapy and chemoradiotherapy.

Discussion

Microvilli control the transport of metabolites and fluids in the polarized epithelial tissues, and their length and density decide the transporting capacity of fluid and metabolites in tissues [1,13]. By analyzing the basal microvilli parameters in 106 pancreatic cancer patients, we found that basal microvilli are common features of the microvasculature in pancreatic cancer. The basal microvilli parameters are equally effective in revealing glucose uptake capacity in pancreatic cancer tissues as PET-CT SUVmax values. They could predict the invasive, metastatic, and proliferative potential of neoplastic cells; prognosis; and therapeutic outcomes of adjuvant therapies.

Genetic and neoplastic cellular alterations are generally used to define the lethality of solid tumors [21]. To the best of our knowledge, basal microvilli are the first tumor-specific microvilli in the tumor microcirculation that indicate the cancer’s lethality. Technically, basal...
microvilli are spatial cellular structures that can be visualized only with a 3D high-resolution scanning method. Thus, our findings imply the potential of 3D high-resolution imaging in visualizing tumor-specific structures, such as basal microvilli, and predicting the pathophysiological characteristics of neoplastic cells in clinical practice. Pathology is still the gold standard for clinical diagnosis and is critical for choosing the treatment regimen. The traditional pathological examination, which relies on the cellular morphology and the spatial arrangement of neoplastic cells, cannot provide precise information about tumor metabolism or the potential

Figure 5. Pancreatic cancer patients with longer and denser basal microvilli are refractory to adjuvant therapy. (A, B) Kaplan–Meier curves for RFS and OS stratified by basal microvilli parameters in the postoperative adjuvant therapy subgroup. Median values of basal microvilli length or density were used as a parameter (length: shorter, <11.17 μm; longer, ≥11.17 μm; density: lower, <6.04/100 μm²; higher, ≥6.04/100 μm²; index: lower, <2.14; higher, ≥2.14). Cox regression was used to assess the hazard ratios; the log-rank test was used to test the P values.
Implications of basal microvilli in pancreatic cancer

metastatic or aggressive nature of neoplastic cells. However, not only are the basal microvilli cancer-specific but they could also provide information about metabolic preferences and the metastatic or invasive and drug-resistant potential of neoplastic cells in vivo.

More than 90% of all pancreatic cancers harbor KRAS mutations [22]. RAS-mutated tumors heavily depend on the Warburg effect – glycolysis to obtain energy for ATP generation [23,24]. The basal microvilli represent systematic changes in the pancreatic microcirculation and are abundant in tumor tissues with elevated glucose uptake. Microvilli of the syncytiotrophoblast in the placenta facilitate the bidirectional exchanges between mother and fetus [13]. In the intestine, microvilli increase the apical surface area by 9.2–15.7 times [25]. In the pancreatic microvasculature, the presence of basal microvilli increases the basal surface area by nearly 0.69–6.07 times. Thus, the microvasculature with basal microvilli might create a ‘super’ microvessel network that is a highly efficient and sustainable shuttle system for transporting glucose and metabolites but not oxygen, perfectly compensating for the hypomicrovascularity in pancreatic cancers. Basal microvilli selectively transport necessities such as nutrients to, or waste from, tumors but not chemotherapeutic drugs. The selective trafficking of basal microvilli might explain our findings that pancreatic cancers with longer and denser basal microvilli are refractory to gemcitabine-based adjuvant therapy and support the notion that basal microvilli might not efficiently deliver gemcitabine to the tumor milieu. Basal microvilli contain caveolae, which are clathrin-independent trafficking vesicles [11,26]. Albumin is reliant on caveolae-mediated endocytosis to be transported or absorbed in pancreatic cancer [27]. Thus, nab-paclitaxel (Abraxane) in pancreatic cancers might cling onto basal microvilli to be transported to cells within the tumor milieu [28].

Our findings also raise several questions concerning the innate characteristics of basal microvilli. Contrary to apical microvilli in other epithelia, basal microvilli appear on the basal surface of endothelial cells in pancreatic cancers. How can these microvessels with basal microvilli traffic nutrients to the tumor milieu? What are the cues for the tumor milieu to induce endothelial cells to form basal microvilli? Currently, high-throughput screening technologies can easily provide candidate cues for basal microvilli. However, a suitable model of basal microvilli in vitro and in vivo is necessary to decode the mechanisms behind the formation of these basal microvilli. Troubling us is the fact that the basal microvilli of human pancreatic cancers decline significantly and diminish quickly when cultured in vitro [12]. This finding suggests that the growth and maintenance of basal microvilli might rely on blood flow. We also discovered basal microvilli in KRAS-mutated autochthonous pancreatic cancers of mice but not in orthotopic pancreatic cancers of mice [11,12,29]. However, the sizes of these tumors in mice are far smaller than those in human cancers. Accordingly, whether autochthonous pancreatic cancer mice are suitable for exploring the regulatory molecules of basal microvilli is questionable.

Collectively, our data showed that the abundance of basal microvilli defines the glucose metabolic capacity and lethal phenotype of pancreatic cancers, indicating that basal microvilli are an anatomical tunnel to deliver drugs or starve neoplastic cells. These results also highlight the emerging role of basal microvilli and suggest an expanding utility in future pathology or clinical practices in pancreatic cancer.

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Author contributions statement

HS and WHL conceptualized the design. HS, XH, WHL, LJ and DLF set up this project. XH and JG collected fresh samples and performed the follow-up for the patients. LXM coordinated the tissue collection. LXM, WHL, XH, JL and DLF provided support for the projects, including from the National Natural Science Foundation (81702304) and SinoGerman Precision Medicine in Pancreatic Cancer (GZ1456). LXM and HS performed all staining, scanning, and measuring of the basal microvilli. HS and XH analyzed the data and wrote the paper.

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SUPPLEMENTARY MATERIAL ONLINE

Figure S1. Basal microvilli patterns

Figure S2. Basal microvilli patterns in pancreatic cancer with an SUVmax of 29.6 on PET-CT and another case with an SUVmax of 4.6 on PET-CT

Figure S3. Longer and denser basal microvilli in pancreatic cancer are related to dismal prognosis (related to Figure 3)

Figure S4. Longer and denser basal microvilli in pancreatic cancer are refractory to adjuvant therapy (related to Figure 4)

Figure S5. Longer and denser basal microvilli in pancreatic cancer are related to dismal prognosis in the no-treatment group (related to Figure 4)

Table S1. Antibodies, equipment, and software list

Table S2. Values of the basal microvilli density, length, and index in 106 pancreatic cancer patients