Cancer-Associated Lymphatic and Venous Vessels in Colonic Carcinomas

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Abstract

Objective: Colonic carcinomas spread to regional lymph nodes and liver. There are cancer-associated lymphatic and venous vessels at the margin of colonic carcinomas, which facilitate spreading carcinoma through lymphatic and venous vessels. This study aimed to examine cancer-associated lymphatic and venous vessels in TNM T1 to T3 carcinomas using lymphatic vessel hyaluronan receptor for lymphatic vessels and von Willebrand factor for venous vessels by immunocytochemical staining.

Materials and Methods: A total of 40 cases of moderately differentiated colonic carcinoma were studied using routinely formalin-fixed and paraffin-embedded sections. The cases consisted of 10 cases of TNM T1, 15 cases each of T2 and T3 cases. Immunocytochemical staining was performed using goat antihuman LYVE-1 for lymphatic vessels and rabbit antihuman von Willebrand factor for venous vessels. Results: In TNM T1 carcinoma, increased, irregular and narrow lymphatic and venous vessels were present in the adjacent normal mucosa to the carcinoma, which penetrated cancerous lesion. There were no tumor emboli in lymphatic and venous vessels. In TNM T2 carcinoma, there were few lymphatic and venous vessels in midst of the carcinoma whereas numerous small lymphatic and venous vessels were present within muscle layers adjacent to the invading carcinoma. Extramural tumor embolus was present in submucosa in one case. In TNM T3 carcinoma, cancer has invaded through the muscle layers where dilated lymphatic and venous vessels were present adjacent to cancerous nests. Tumor emboli were identified in two cases by immunocytochemical staining. Conclusion: The current study showed cancer-associated lymphatic and venous vessels at the interface in TNM T1 carcinoma to dilated intramuscular lymphatic and venous vessels adjacent to invading cancerous nests in TNM T3 carcinoma, and supports cancerous cells spread via lymphatic and venous vessels through muscle layers to subserosa as supported by tumor emboli in the lymphovascular system.

Keywords
Colonic Carcinoma, Factor-8, Lymphatic Vessels, LYVE-1, Venous Vessels

1. Introduction

The most common sites of metastasis of colonic carcinoma are regional lymph nodes and liver [1]. Lymph node involvement and number of metastatic lymph nodes are considered to be an important clinicopathological factor in colorectal carcinoma [1]-[5]. Since cancer cells metastasize to regional lymph nodes through lymphatic vessels and to liver through venous vessels, lymphatic and venous vessel tumor invasion is the histopathological risk factor for colorectal carcinoma [6]. It is not easy to definitely identify and evaluate lymphatic vessel invasion by routine H. and E. sections, and immunocytochemical identification for the lymphatic vessel is a currently available tool for identifying lymphatic vessels [7]. There is an agreement on the evidence that more lymphatic and venous vessels are present at the margins of colonic carcinoma, suggesting that colonic carcinoma spreads to regional lymph nodes through lymphatic vessels [1]-[5] and to liver by hematogenous spread [8] [9]. But cancer-associated lymphangiogenesis has been a controversial issue [3]-[5] partly due to a lack of specific and reliable immunocytochemical markers for lymphatic endothelium [10]-[12] and a lack of universally accepted methodology to identify and evaluate lymphatic vessels in surgically resected colonic cancer specimens.

By immunocytochemical staining using lymphatic vessel endothelial hyaluronan receptor-1 (LYVE-1) for lymphatic vessels and von Willebrand factor (factor-8, F-8) for venous vessels, lymphatic and venous vessels were concomitantly studied in surgically resected colonic cancer specimens since this immunocytochemical staining was previously used with colonic specimens harboring polyps and adenomas [13]. Recently, immunocytochemical staining for lymphatic vessels has been performed using Prox1, podoplanin, D2-40 and LYVE-1, the latter has been widely used as a reliable marker in the previous studies [7] [10]-[13].

2. Materials and Methods

All cases of moderately differentiated colonic carcinoma including the adjacent normal colon were from the University of Kansas Medical Center, collected between 1999 and 2000, and consisted of 10 cases TNM T1, 15 cases each of T2 and T3 stages of carcinomas with a total of 40 cases [14]. Two tissue blocks including the interface between the cancerous lesion and normal mucosa were studied for T1 cases and two sections of the interface containing the entire colonic wall thickness were studied for T2 and T3 cases. The surgically removed specimens were routinely fixed in buffered formalin and were embedded in paraffin. Deparaffinized sections were treated with antigen retrieval procedure using citrate buffer pH 6.2. Immunocytochemical staining was performed for LYVE-1 using goat antihuman LYVE-1 (R & D System, Minneapolis, MN) at 1:100 dilution and F-8 immunocytochemical staining was performed using rabbit human F-8 (Dako System, Carpenteria, CA) at 1:100 dilution as reported before [13]. At least two sections from each tissue block at the two different levels were immunostained for LYVE-1 and F-8. Only the sections, which were adequately immunostained for both LYVE-1 and F-8, were included in this study.

3. Results

Normal colon: Abundant LYVE-1 positive lymphatic vessels were slender, irregularly shaped channels in submucosa including muscularis mucosa, which abruptly changed the directions and consistently extended into the base and deeper end of colonic crypt while extending into submucosa vertically through the horizontal muscularis mucosa and also numerously into lymphoid follicles, when the latter were present (Figure 1(A)). Less abundant F-8 positive venous vessels appeared of more roundish shape with plump lumens, and extended sparingly into both submucosa and the entire thickness of lamina propria onto the mucosal surface and into lymphoid follicle (Figure 1(B)). Both lymphatic and venous vessels penetrated vertically through the inner muscular layer whereas the two vessels spread horizontally in the outer longitudinal muscle layer in the tangentially cut sections against the mucosal folds (Figure 1(C) and Figure 1(D)). Both intramuscular lymphatic and venous vessels were smaller and slender than those in the submucosa (Figures 1(A)-(D)). In the parallel cut sections along the
Figure 1. Normal colon. Lymphatic vessels were abundant in submucosa, penetrating through muscularis mucosa and into the base and deep-end of colonic crypt and numerously into submucosal lymphoid follicle (A). Venous vessels were relatively less numerous than lymphatic vessels in submucosa and were round with plump lumens, penetrating through muscularis mucosa sparsingly into the full thickness of lamina propria onto the mucosal surface (B). Numerous small lymphatic and venous vessels penetrated vertically through the inner circular muscle layer whereas the both vessels spread horizontally in the outer longitudinal muscle layer in the tangentially cut sections against the mucosal folds (C) and (D). I: Inner muscle layer, O: Outer muscle layer, L: Lymphoid follicle. (A) and (C): LYVE-1; (B) and (D): F-8 immunostaining.

Colonic carcinoma, TNM T1: There was thin fibrous stroma between back-to-back cancerous glands, between which were a few lymphatic vessels in contrast to small but more abundant venous vessels (Figure 2(A) and Figure 2(B)). In the interface between the cancerous and normal mucosa, there were more lymphatic vessels and much more venous vessels in the adjacent normal mucosa, some of which were continuous to the cancerous mucosa (Figure 2(A) and Figure 2(B)). At the surface of cancerous mucosa, there were focally abundant small lymphatic and venous vessels in fine fibrous stroma (Figure 2(C) and Figure 2(D)). Some of the vessels in the interface were immunostained concomitantly for both LYVE-1 and F-8 (Figure 2(A) and Figure 2(B)). Mid cancerous mucosa revealed less lymphatic and venous vessels than the distant normal mucosa. The submucosa adjacent to the cancerous lesion revealed numerous slender lymphatic vessels and larger round venous vessels,
Figure 2. Interface of TNM T1 adenocarcinoma. In mucosal interface between carcinoma and normal mucosa, there were moderately increased lymphatic vessels and much more increased venous vessels in the adjacent normal mucosa (A) and (B). Some small vessels (n) were positively immunostained for both LYVE-1 and F-8 (n). At the cancerous mucosal surface, there were focally increased small lymphatic vessels and more increased venous vessels in the fine fibrous stroma (C) and (D). L: Lymphoid follicle, n: vessels stained for both LYVE-1 and F-8. (A) LYVE-1, (B) F-8 immunostaining.

and there were no tumor emboli in lymphatic and venous vessels.

Colonic carcinoma, TNM T2: In submucosa, there were abundant lymphatic and venous vessels with tumor emboli identified in 1 of each 15 cases of lymphatic (Figure 3(A)) and venous vessels (Figure 3(D)). Adjacent to the invading tumor within the muscle layers, there were numerous small lymphatic and larger venous vessels, which have vertically penetrated through the muscle layer (Figure 3(B) and Figure 3(C)). The majority of cancerous nests in the submucosa were surrounded by an empty halo, not representing lymphatic or venous vessels but appeared to be a contraction artifact by formalin fixation and paraffin embedding, supported by the negative immunostaining for both LYVE-1 and F-8 (Figure 3(A) and Figure 3(D)).

Clonic carcinoma, TNM T3: In muscle layers adjacent to the invading carcinoma, there were abundant small lymphatic and more abundant small venous vessels (Figure 4(A) and Figure 4(B)). Cancerous nests in the muscular layers were surrounded by small lymphatic vessels and abundant small venous vessels, some of the latter penetrated into the cancerous nests (Figure 4(C) and Figure 4(D)). In two cases, there were extramural tumor emboli in lymphatic and venous vessels out of 15 cases. In the subserosa, there were increased small lymphatic vessels and numerous, round venous vessels.

4. Discussion

Colonic cancer spreads predominantly to regional lymph nodes through lymphatic vessels although little is known about cancer-associated lymphangiogenesis [15] [16]. Much of the difficulty in analyzing this lymphangiogenesis is due to a lack of specific lymphatic vessel markers to differentially identify lymphatic and venous vessels by immunocytochemical staining [15]-[17]. Several lymphatic endothelial markers are now available, including LYVE-1, Prox1, podoplanin and D2-40, in which CD 34 is expressed by LYVE-1+/podoplanin+/
Figure 3. Submucosa and muscle layers of TNM T2 adenocarcinoma. In the submucosa adjacent to the invading cancerous nests, there were abundant slender lymphatic vessels (A) and larger and plump venous vessels (B). There were tumor emboli in submucosal lymphatic vessels (A) and venous vessels (D) whereas the majority of cancer nests in submucosa were surrounded by empty halos, not representing lymphatic or venous vessels but appeared to be a contraction artifact by formalin-fixation and paraffin-embedding (A), (C) and (D). (C) was from the section cut along the mucosal folds, showing horizontal inner muscle layer. a: artery, E: tumor embolus, I: Inner muscle layer, O: Outer muscle layer. (A) and (C) LYVE-1, (B) and (D) F-8 immunostaining.

Prox1+ cancer-associated lymphatic cells in colon, breast, lung and skin tumors whereas lymphatic endothelial cell in all normal organs does not express CD 34 by immunofluorescence study on unfixed frozen sections [17].

Matsumoto et al. [18], Kaneko et al. [19] and Liang et al. [20] used podoplanin for “hot spots” and Zimmerman et al. [15], Gao et al. [21] and Longatto-Fielo et al. [22] used monoclonal D2-40 for lymphatic vessel microanalysis in peritumoral tissue of colonic carcinoma and came to the same conclusion that there was a correlation between high lymphatic vessel microdensity and positive lymphatic vessel invasion/lymph node metastasis although these authors did not mention the correlation between lymphatic vessel microdensity and lymphovascular tumor invasion [18]-[22]. Using LYVE-1 immunostaining, Minsky et al. initially studied 462 cases of colonic carcinoma and found 61 cases (13%) showing lymphovascular invasion and had shown that lymphovascular invasion is an independent prognostic factor [23].

Prognosis of colonic carcinoma depends on vertical invasion of cancer cells according to TNM stages [14], and vertical lymphangiogenesis though the muscle layers to the deeper margins may facilitate lymphatic spread
Figure 4. Deep muscle layers and subserosa of TNM T3 adenocarcinoma. There were massive cancerous masses in the muscle layers, which contained few lymphatic and venous vessels whereas there were numerous small lymphatic and venous vessels at the margins of the invading cancer (A) and (B). Adjacent to the invading cancerous nests through muscular layers, small lymphatic and round, numerous venous vessels were noted in the muscular layers (C) and (D), some of which were continuous into the cancerous nests (D). There were more round venous vessels surrounded the cancerous nests than smaller lymphatic vessels in the subserosa (C) and (D). (A)-(D) were from the sections cut along the mucosal folds, showing horizontal inner muscle layer and vertical outer muscle layer. a: artery, l: Inner muscle layer, L: Lymphoid follicle, O: Outer muscle layer, s: subserosa, t: tumor nests, v: vein. (A) and (C) LYVE-1, (B) and (D) F-8 immunostaining.

of the carcinoma. Both lymphatic and venous vessels penetrate vertically and horizontally through thicker circular inner muscle bundles and through the thinner outer longitudinal muscle layer (Figure 1 and Figure 2). Both lymphatic and venous vessels in the muscle layers are generally smaller than those in submucosa and subserosa.

The current study showed both horizontal and vertical lymphatic and venous vessel proliferation at the adjacent margins of the mucosal interface and submucosa (Figure 3(A) and Figure 3(D)) and the normal submucosa adjacent to the cancer contained abundant larger lymphatic and venous vessels. Some of the extramural vessels even contained tumor emboli as a proof for cancerous cells transported through the lymphovascular system (Figure 3(A) and Figure 3(D)). This lymphangiogenesis appeared to be cancer-associated as abundant lymphatic vessels were noted adjacent to the tumors and the invading cancerous nests (Figure 3, Figure 4). In TIM T3 carcinomas, increased, small intramuscular lymphatic vessels were present (Figure 4(C) and Figure 4(D)).
These numerous lymphatic and venous vessels were noted adjacent to the invading carcinoma in TNM T2 and T3 cases (Figure 3 and Figure 4) in contrast to scanty lymphatic and venous vessels in mid portion of TNM T1 carcinomas (Figure 2). Therefore, cancer-associated lymphangiogenesis and angiogenesis were predominantly observed adjacent to the invading T2 and T3 carcinoma (Figure 3 and Figure 4).

Some cancer-associated small vessels were positive for both LYVE-1 and F-8 (Figure 2(A) and Figure 2(B)), suggesting newly formed cancer-associated lymphangiogenesis from common stem cells for lymphatic and venous vessels [10]-[13]. We had also used D2-40 (Dako System) for immunostaining lymphatic vessels for lymphatic vessels, which revealed no lymphatic vessels in the mid cancerous tissue similar to those observed by LYVE-1 immunostaining (Unpublished data).

Using routinely processed formalin-fixed and paraffin-embedded sections, not all lymphatic vessels were consistently immunostained for lymphatic and venous vessels in our experience as also reported by others [13] [24] [25], most likely due to non-optimal fixation and/or poor tissue preservation for the epitopes. Furthermore, even in the well formalin-fixed and paraffin-embedded sections, lymphatic vessels were not evenly immunostained but were occasionally focally and sporadically immunostained within the same cases or even in the same microscopic slides, likely due to uneven preservation for the epitopes. It is apparent that formalin-fixation and paraffin-embedding masked or destroyed some parts of the LYVE-1 and F-8 epitopes. Fiedler et al. used frozen sections for immunofluorescence staining for multiple lymphatic endothelial markers [17] but this procedure is cumbersome and too labor intensive for a routine pathology laboratory. Complex histopathological and functional interactions between cancerous cells and lymphangiogenesis in the cancer may definitely play a pivotal role in the commonly metastasizing tumor like colonic carcinoma [26].

This current study has shown that there were abundant small extramural lymphatic and more abundant venous vessels in T1 carcinomas and, there were more abundant intramural lymphatic and venous vessels once cancer has invaded into muscle layers in TNM T2 carcinoma, and further more abundant intramural lymphatic and venous vessels appeared when the cancer invaded through the muscle layers to subserosa in TNM T3 cases. These cancer-associated lymphatic and venous vessels appeared concomitantly and may provide an easy access for cancer cells to regional lymph nodes and beyond, including hematogenous spread to liver. Lymphovascular invasion identified by tumor emboli was 0% to 15% in our cases. Harris et al. assembled a group of six GI pathologists to test an interobserver variability by examining 50 cases of moderately-differentiated T1 to T3 colorectal carcinoma and the tally was: 7%, 24% and 31% for lymphatic invasion and 0%, 10% and 23% for venous vessels for T1, T2 and T3 the carcinomas, respectively [27]. There was substantial interobserver variability in diagnosing lymphovascular invasion among the six pathologists due to the different individual criteria for lymphovascular invasion [27]. Immunocytochemical staining for D2-40 for lymphatic vessels and CD 31 for venous vessels did not improve for identifying lymphovascular invasion using the currently available antibodies [27]. This study highlights the need for criteria in evaluation of lymphovascular invasion [27]. Thus, a reported incidence of lymphovascular invasion for colonic carcinomas was quite variable from 0% to 31% using H. and E. sections and immunostained sections for D2-40 and CD 31 [27]-[29]. For a better immunocytochemical study, better antibodies with a higher specificity and sensibility than the currently available sources are warranted. Nevertheless, lymphovascular invasion in colonic carcinoma still remains as a strong stage-independent prognostic marker [30] [31] and this immunocytochemical study has to be pursued in the future.

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