Tumor cell invasion from the marginal sinus into extranodal veins during early-stage lymph node metastasis can be a starting point for hematogenous metastasis

Tetsuya Kodama1,2, Shiro Mori1,2,3, Masato Nose4

1Graduate School of Biomedical Engineering, Tohoku University, Sendai 980-8575, Japan.
2Biomedical Engineering Cancer Research Center, Graduate School of Biomedical Engineering, Tohoku University, Sendai 980-8575, Japan.
3Department of Oral and Maxillofacial Surgery, Tohoku University Hospital, Sendai 980-8575, Japan.
4Institute for Promotion of Advanced Science and Technology, Ehime University, Matsuyama 790-8577, Japan.

Correspondence to: Prof. Tetsuya Kodama, Graduate School of Biomedical Engineering, Tohoku University, 4-1 Seiryo, Aoba, Sendai 980-8575, Japan. E-mail: kodama@tohoku.ac.jp

Abstract

Aim: To investigate whether tumor cells in a lymph node (LN) can invade from the marginal sinus into extranodal veins via vessel branches that communicate with intranodal veins and whether this can be a starting point for hematogenous metastasis at the early stage of LN metastasis.

Methods: Vascular and lymphatic networks of LNs in MXH10/Mo-lpr/lpr mice were investigated using three-dimensional micro-computed tomography and histological methods. Flow in the blood vessel networks of LNs was investigated by fluorescence microscopy. Tumor cells were injected into the subiliac LNs of MXH10/Mo-lpr/lpr mice to induce metastasis to the proper axillary LNs. Tumor development in the proper axillary LN was detected using an in vivo bioluminescence imaging system. A two-dimensional image of the proper axillary LN microvasculature was reconstructed using a contrast-enhanced high-frequency ultrasound system.

Results: Extranodal veins communicated with intranodal veins via branches that penetrated the capsule, and blood flowed from intranodal veins to extranodal veins. Tumor cells that had metastasized to the marginal sinus invaded these communicating veins to develop hematogenous metastases.
Conclusion: Metastatic LNs that would be considered by clinical imaging to be stage NO can be a starting point for hematogenous metastasis. The study findings highlight the need for the development of novel techniques for the diagnosis and treatment of early-stage LN metastasis, i.e., when standard diagnostic imaging might incorrectly classify the LN as stage NO.

Keywords: Lymph node, metastasis, NO, lymph node-mediated hematogenous metastasis

INTRODUCTION

Tumor cells reach the marginal sinus of a sentinel lymph node (LN) via afferent lymphatic vessels, after which they proliferate along the lymphoid sinus, invade the cortex and reach the medulla. The abundant vascular network in a LN allows tumor cells to grow without the induction of tumor neovasculature. Since tumor cells growing in a LN can infiltrate both the lymphatic channel and the vascular network, a sentinel LN can be the origin of lymphatic metastasis to downstream LNs as well as hematogenous metastasis. It has been suggested that high endothelial venules (HEVs) may be involved in the mechanisms underlying systemic metastasis, but the details remain unknown. Clinically, a LN is judged as positive for metastasis (> N1) if tumor invasion is detected by diagnostic imaging or aspiration cytology. However, since a LN can be erroneously classified as stage N0 during the early stages of tumor invasion, a false-No LN can potentially be a source of systemic metastasis. For example, in the NSABP-32 trial, patients with breast cancer judged incorrectly to be stage No had no difference in overall survival, disease-free survival and distant disease-free interval to patients judged to be stage No. In other words, tumor cells may have undergone systemic metastasis at a stage when LNs were incorrectly classified as N0.

Recently, we demonstrated that a fluorescent dye injected locally into a LN flowed into both the efferent lymphatic vessel and extranodal veins and that intranodal and extranodal veins communicated via branches that passed through the capsule, a feature not described in conventional textbooks of anatomy. Thus, tumor cells can undergo both lymphatic and hematogenous metastasis. Based on these results, we proposed a theory of LN-mediated hematogenous metastasis, whereby LNs can be the origin of systemic metastasis.

In this study, we used a mouse model in which metastasis to the proper axillary LN (PALN) was induced by the inoculation of tumor cells into the subiliac LN (SiLN). We found that during early-stage PALN metastasis (confirmed by pathological imaging), the invasion of tumor cells from the marginal sinus into intranodal veins and then extranodal veins may be a first step in the mechanism of hematogenous metastasis from a LN.

METHODS

Experiments were carried out in accordance with published guidelines and approved by the Institutional Animal Care and Use Committee of Tohoku University.

Mice

MXH10/Mo-lpr/lpr (MXH10/Mo/lpr) mice (12-17 weeks old), which are a congenic strain of MRL/Mp-lpr/lpr and C3H/HeJ-lpr/lpr mice, were bred under specific pathogen-free conditions in the Animal Research Institute, Graduate School of Medicine, Tohoku University, Sendai, Japan. The LNs enlarge to about 10 mm in diameter at 12 weeks of age due to invasion by lpr-T (CD4 CD8 B220 Thy) cells. The anatomical locations and nomenclatures of murine LNs have often been ignored or assigned incorrectly; in this study, we used the term “subiliac LN” instead of “inguinal LN”.

Micro-computed tomography imaging

Specimens were analyzed using high-resolution micro-computed tomography (micro-CT) scanning (scanXmate/E090, Comscan Tecnio). Barium contrast agent (mean size, 935.7 nm) was prepared as
previously described\textsuperscript{[1]}\textsuperscript{[12]}. With the mouse under deep general anesthesia, 0.1 mL heparin (Novo-Heparin, 1000 units/mL, Mochida Pharmaceutical) was administered intravenously and 0.05 mL papaverine hydrochloride (40 mg/mL, Nichi-Iko) was given subcutaneously. Ten min later, a syringe pump (Legato100, KD Scientific) was used to infuse 4 mL of saline (18 mL/h) into the left ventricle through a thoracotomy. After cutting of the caudal vena cava and draining of the blood from the body, 4 mL of contrast medium was injected into the left ventricle (18 mL/h) via a T-shaped stopcock. Following perfusion with contrast medium, the LNs were dissected and kept at 4°C for >2 h to allow fixation to occur. The samples were placed on the stage of a micro-CT scanner (the gutta percha was used as a landmark for positioning) and scanned (angiography) at resolutions of 5-30 μm and a slice thickness of 100 μm. Acquired slice data were rendered as 3D images using a 3D analysis suite (Amira, Maxnet).

Cell culture
KM-Luc/GFP cells (mouse malignant fibrous histiocytoma-like cells derived from an MRL/Mp-gld/gld mouse expressing a fusion of the luciferase and enhanced-green fluorescent protein genes) were cultured as previously described\textsuperscript{[14]}. FM3A-Luc cells (C3H/He mouse mammary carcinoma cells expressing the luciferase gene\textsuperscript{[16]}) were maintained in RPMI-1640 medium supplemented with 10% fetal bovine serum, 1% L-glutamine-penicillin-streptomycin and 1 mg/mL G418 (Sigma-Aldrich). Both cell types had an H-2\textsuperscript{k} haplotype, which is the same as that of MXH10/Mo/lpr mice, and expressed vascular endothelial growth factor (VEGF)-A and VEGF-B but not VEGF-C; KM-Luc/GFP but not FM3A-Luc cells showed slight VEGF-D expression\textsuperscript{[15]}. The relative growth rates of KM-Luc/GFP and FM3A-Luc cells were 3.8/day and 1.1/day, respectively\textsuperscript{[8]}. Cell lines were incubated (37°C, 5% CO\textsubscript{2}/95% O\textsubscript{2}) until 80% confluence was achieved. Lack of Mycoplasma contamination was confirmed on the inoculation day (MycoAlert Mycoplasma Detection Kit; Lonza Rockland).

NBD-liposomes
NBD-liposomes were synthesized from 1,2-distearoyl-sn-glycero-3-phosphatidylcholine (DSPC; MC8080, NOF Co.), 1,2-distearoyl-sn-glycerol-3-phosphatidylethanolamine-methoxy- polyethyleneglycol (DSPE-PEG[2000-OMe]; DSPE-020CN, NOF Co.), and 1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine-N-(7-nitro-2-1,3-benzoxadiazol-4-yl) (NBD-DPPE; FE6060, NOF Co.)\textsuperscript{[9,16]}. The size and zeta potential of the NBD-liposomes was 642 nm and -1.5 mV, respectively, as measured using a particle size and zeta potential analyzer (ELSZ-2; Otsuka Electronics).

Visualization of venous flows internal and external to a LN
Under deep general anesthesia, an arc-shaped incision was made in the abdominal skin of a mouse (\(n = 1\), 12 weeks old) from the subiliac region to the proper axillary region, and 100 μL of 0.01 μmol/L NBD-liposomes was injected into the tail vein (100 μL/min, 60 s). Fluid flow in the veins was visualized using a fluorescence stereomicroscope (M165-FC; fluorescence filter: GFP2; excitation: 460-500 nm; emission: > 510 nm; Leica) connected to a high-speed camera (10 frames/s; CoolSNAP HQ2; Photometrics)\textsuperscript{[8]}.

Induction of metastasis to the PALN by injection of tumor cells into the SiLN
KM-Luc/GFP (1.5 × 10\textsuperscript{6} cells/mL) or FM3A-Luc (3.3 × 10\textsuperscript{5} cells/mL) cells (passaged three times) were suspended in a mixture of 20 μL phosphate-buffered saline (PBS) and 40 μL of 400 mg/mL Matrigel (Collaborative Biomedical Products). The prepared cells were injected into the center of the SiLN of a mouse (aged 14-16 weeks) using a 27G needle, which was maintained in the same position for 5 min to allow solidification of the Matrigel. Tumor development in the SiLN and metastasis to the PALN (the rates of which depended on the tumor cell type) were detected using an in vivo bioluminescence imaging system (IVIS; Xenogen) at 4 and 7 days post-inoculation of KM-Luc/GFP cells and at 6, 13, 20 and 27 days post-inoculation of FM3A-Luc cells. Injection of tumor cells into the SiLN induces metastasis to the PALN via lymphatic vessels as well as hematogenous metastasis via the thoracoepigastric vein (TEV)\textsuperscript{[9,17]}; the cell number is highest in the SiLN and second highest in the PALN\textsuperscript{[14]}.
Contrast-enhanced ultrasound imaging and spatiotemporal analysis of pixel intensity variations

A 2D image of the PALN microvasculature was reconstructed on days 6 and 9 post-inoculation of KM-Luc/GFP cells and on days 8, 15, 22 and 29 post-inoculation of FM3A-Luc cells using a contrast-enhanced high-frequency ultrasound (CE-HFUS) system with a 37.5-MHz transducer (RMV-710B, VisualSonics). Each transducer was fixed to a 3D stage control system (Mark-204-MS; Sigma Koki). Contrast images (slice thickness, 100 µm) were captured before and 60 s after an intravenous bolus injection of 100 µL microbubbles (Sonazoid, Daiichi Sankyo) into the tail vein. During imaging, the mouse was positioned on a heated stage and anesthetized with 2% isoflurane in oxygen. Acquired images were analyzed using ultrasound contrast agent-detecting software to measure temporal changes in the PALN microvessel density.

Histological analysis

PALNs and SiLNs were harvested at the indicated time points, fixed in 10% formaldehyde in PBS for 3 days, placed on a shaker for 1 day at room temperature and then stored at 4 °C for 2 days. Next, the samples were dehydrated in 100% ethanol, placed into a tissue processor, embedded in paraffin and cut into 3-µm serial sections using a fully-automated tissue-sectioning device (AS-400, Kurano). Samples were stained with hematoxylin and eosin (HE) or immunostained for detection of CD31-positive vascular endothelial cells.

Statistical analysis

Data are presented as mean ± SD or mean ± SEM. Differences between groups were determined by one-way analysis of variance followed by Tukey’s test or an unpaired t-test (GraphPad Prism 6). *P < 0.05 was considered to be statistically significant.

RESULTS

Connection of the TEV to SiLN blood vessels

First, we examined the anatomical positions of the PALN, SiLN, accessory axillary LN (AALN) and TEV. An arc-shaped incision was made in the abdominal skin of a mouse from the subiliac region to the proper axillary region. The axillary area contains two LNs, the PALN and AALN. The SiLN and AALN lie upstream of the PALN in the lymphatic network. The TEV, which connects the subclavian vein and inferior vena cava, runs adjacent to the PALN and SiLN along the lymphatic vessels (not visualized in Figure 1A) between these LNs. The TEV receives venous blood from the PALN and SiLN via small branches. There were many vessels on the LN that connected to intranodal vessels. The hilum of the PALN and SiLN was located behind the image.

A series of 3D micro-CT images, rotated every 60°, revealed a complex vascular network in the PALN, with many small branches penetrating the LN capsule and connecting the TEV to intranodal blood vessels. There were no similar networks on the reverse side. Immunolabeling of CD31 showed that the TEV ran along and penetrated the capsule of the SiLN along the lymphatic vessels (Figure 1C) between these LNs. The TEV receives venous blood from the PALN and SiLN via small branches. There were many vessels on the LN that connected to intranodal vessels. The hilum of the PALN and SiLN was located behind the image.

Flow dynamics between the PALN blood vessel network and TEV

The flow dynamics between the PALN blood vessel network and TEV were visualized under a fluorescence stereomicroscope after intravenous injection of NBD-liposomes. Consistent with the results shown in Figure 1, the TEV communicated with intranodal blood vessels via many small branches. In the region excluding the TEV, a large vein running along the PALN made connections with the intranodal veins, and the combined blood flow drained into the TEV. In the region including the TEV, the venous network inside the LN connected to the TEV.
Figure 1. Vascular and lymphatic networks of lymph nodes (LNs). A: Vascular networks in LNs: (I) a macroscopic view of a 14-week-old mouse (right side) indicating the anatomical positions of the proper axillary LN (PALN), subiliac LN (SiLN), accessory axillary LN (AALN) and thoracoepigastric vein (TEV). The axillary area contains two LNs, the PALN and AALN\[^{16}\]. The SiLN and AALN are upstream of the PALN in the lymphatic network. The TEV, which connects the subclavian vein (SV) and inferior vena cava (IVC), runs adjacent to the SiLN and PALN and along the lymphatic vessels between them (not visualized). The TEV receives venous blood from the SiLN and PALN via small branches; (II) blood vessels running on the PALN in a 17-week-old mouse. There were many vessels on the LN surface. The hilum was behind the image; (III) blood vessels running on the SiLN in a 17-week-old mouse. There were many vessels on the LN surface. The hilum was behind the image: B: three-dimensional micro-computed tomography (micro-CT) images showing the surface and internal vascular structure of the PALN in a 14-week-old mouse. The series of images shows the LN rotated by 60° each time. The TEV communicated with intranodal blood vessels via many small branches that penetrated the LN capsule. There were no similar networks on the reverse side. Scale: 2 mm. g: gutta-percha; C: images immunostained for CD31 showing the connections between the TEV and SiLN blood vessels in a 16-week-old mouse. Images I-IV (middle and right) are magnified views of the corresponding regions highlighted in the left-most image. The TEV ran along the capsule (image I) and connected with intranodal veins penetrating through the marginal sinus (image II). It is notable that the marginal sinus was extremely close to the intranodal vein. The branches of TEV ran in the cortex under the marginal sinus (image III) and branched into two vessels in the cortex (image IV). RBC: red blood cell; Cap: capsule; Mgs: marginal sinus; Cor: cortex; Paracor: paracortex
from the TEV enters the subclavian vein; since the venous branches from the intranodal veins connect to the TEV, their blood is returned to the venous circulation.

**Metastatic SiLN at the false N0 stage**

Next, we explored the interaction of tumor cells with intranodal vessels during the early stage of metastasis. Two different tumor cell types were used: KM-Luc/GFP [Figure 3A, C, E] and FM3A-Luc [Figure 3B, D, F]. The injection of tumor cells into the SiLN induced metastasis in the PALN. Luciferase activity in the SiLN and PALN increased with tumor progression for both cell lines (Figure 3A and C for KM-Luc/GFP, Figure 3B and D for FM3A-Luc). Metastasis was detected on day 7 after inoculation of KM-Luc/GFP cells [Figure 3A] and day 27 after inoculation of FM3A-Luc cells [Figure 3B]. Subsequently, we investigated flow dynamics in and around the metastatic PALN using CE-HFUS [Figure 3E and F]. Microbubbles flowing in the TEV were visualized in experiments using KM-Luc/GFP cells [Figure 3E and Video 3], while confluence of the TEV with other vessels was visualized in experiments using FM3A-Luc cells [Figure 3F and Video 4]. Our previous studies demonstrated no significant change in the volume of the metastatic PALN up to day 7 for KM-Luc cells [20] and day 22 for FM3A-Luc cells [21]. Thus, in a clinical setting, the metastatic PALN in these experiments would have been classified as stage N0 by diagnostic ultrasound.

**Interaction of tumor cells with surrounding blood vessels in a LN at the false N0 stage**

Histological techniques were used to investigate the interaction of tumor cells in the PALN (at the false N0 stage) with the surrounding blood vessels at day 6 for KM-Luc/GFP cells [Figure 4A and B] and day 8 for FM3A-Luc cells [Figure 4C-E]. The PALNs were removed after the micro-CT imaging experiments had been completed so that the blood vessels were filled with contrast agent. In experiments using KM-Luc/GFP cells, serial sections stained with HE [Figure 4A and B, Supplementary Figure 1] or immunostained for CD31 [Supplementary Figure 1] showed an afferent lymphatic vessel entering the LN. Tumor cells from the afferent lymphatic vessel had invaded the marginal sinus as well as vessels that were filled with contrast agent. In experiments using FM3A-Luc cells [Figure 4C and D], metastasized tumor cells were detected in the marginal sinus of the PALN. Importantly, tumor cells that had developed in the marginal sinus had...
Figure 3. Induction of metastasis of KM-Luc/GFP cells or FM3A-Luc cells to the proper axillary lymph node (PALN) after inoculation of the cells into the subiliac lymph node (SiLN). KM-Luc/GFP cells (A, C, E); FM3A-Luc cells (B, D, F). A: Progression of KM-Luc/GFP cell metastasis from the SiLN to the PALN, as detected from measurements of luciferase activity; B: progression of FM3A-Luc cell metastasis from the SiLN to the PALN, as detected from measurements of luciferase activity; C: quantification of luciferase activity in the SiLN and PALN after inoculation of KM-Luc/GFP cells into the SiLN (SiLN, n = 5 at day 4 and n = 3 at day 7; PALN, n = 4 at day 4 and n = 4 at day 7). ****P < 0.0001, unpaired t-test. The bars show mean ± SEM values; D: quantification of luciferase activity in the SiLN and PALN after inoculation of KM-Luc/GFP cells into the SiLN (SiLN, n = 14 at day 6, n = 10 at day 13, n = 7 at day 20 and n = 3 at day 27; PALN, n = 14 at day 6, n = 10 at day 13, n = 7 at day 20 and n = 3 at day 27). ***P = 0.0001, ****P < 0.0001, one-way ANOVA and Tukey’s test. The bars show mean ± SEM values; E: representative images of the PALN at day 6, obtained using contrast-enhanced high-frequency ultrasound (CE-HFUS; n = 5; Video 3), in a mouse inoculated with KM-Luc/GFP cells. The CE-HFUS image reveals microbubbles flowing in the vessel in the PALN; F: representative images of the PALN at day 8, obtained using CE-HFUS (n = 5; Video 4), in a mouse inoculated with FM3A-Luc cells. The CE-HFUS image shows microbubbles flowing in the vessel in the PALN. NS: not significant.
Figure 4. Invasion of tumor cells into a subcapsular vein in the proper axillary lymph node (PALN). A, B: Sections of the PALN at day 6 after inoculation of KM-Luc/GFP cells into the subiliac lymph node (SiLN). The sections were stained with hematoxylin-eosin (HE); the interval between each section was 12 μm. Tumor cells (T) that had entered from the afferent lymphatic vessel (ALV) had invaded into the vein in the marginal sinus (Mgs) of the PALN. Scale bar: 100 μm. Cor: cortex; CA: contrast agent; C-F: PALN at day 8 after inoculation of FM3A-Luc cells into the SiLN [C: tumor cells localized in the marginal sinus of the PALN. HE staining. Scale bar: 200 μm; D: higher magnification of the rectangular region in (C). The marginal sinus was filled with tumor cells in a botryoidal configuration that projected toward the vascular wall. HE staining. Scale bar: 50 μm. E: higher magnification of the rectangular region in (D). The tumor cells had disrupted the littoral cell layer of the marginal sinus and the vascular wall and had invaded into the vascular lumen (arrow). HE staining. Scale bar: 20 μm; F: illustration of (D) showing the invasion of metastatic tumor cells from the marginal sinus into the subcapsular vein. Cap: capsule; Cor: cortex; Mgs: marginal sinus (subcapsular lymphatic sinus).]
entered the region close to the subcapsular vein and invaded the vessel [Figure 4E]. The schematic diagram in Figure 4F illustrates the internal features of the marginal sinus with metastatic tumor cells invading the subcapsular vein.

DISCUSSION

The present study demonstrates that tumor cells in the marginal sinus of a LN can invade extranodal veins via branches that communicate with intranodal veins. We suggest that this may be the first step in hematogenous metastasis from LNs. Importantly, this form of metastasis occurs before tumor cells have interfered with well-developed vascular networks and HEVs within the LN\(^\text{[5,6]}\). Veins are ubiquitously present on the surface layer of the LN [Figure 1]\(^\text{[22]}\) and communicate with intranodal veins, allowing blood to flow to the systemic circulation from intranodal vessels. This anatomical arrangement and flow characteristics have not been described previously. Kelch et al.\(^\text{[2]}\) surgically excised individual mesenteric LNs from the mouse and clarified the topology of the entire intranodal vascular network using an automated confocal imaging system and custom-written software. However, unlike the present study, their analysis was limited to the vascular network within the LN. This study utilized malignant fibrous histiocytoma-like KM-Luc/GFP cells and breast cancer FM3A-Luc cells, both of which are syngeneic to the recipient mice. The invasion of tumor cells from the afferent lymphatic vessel into the intranodal veins was confirmed for both cell types [Figure 4, Supplementary Figure 1]. This implies that LN-mediated hematogenous metastasis may be a mechanism relevant to a wide variety of cancer types.

Lymphadenopathy in MXH10/Mo/lpr mice is due to the lpr gene and is characterized by the accumulation of lpr-T cells in the paracortical region. The metastatic lesions shown in Figure 4 are located in the marginal sinuses of the LNs. Thus, it is unlikely that there is a direct relationship between abnormal lymphocyte proliferation in the LN parenchyma and tumor cell proliferation and vascular invasion.

MXH10/Mo/lpr mice do not express the fas gene involved in apoptosis, since the lpr gene is a fas-deletion mutant gene. Thus, the immune system in MXH10/Mo/lpr mice is functional except for the signaling pathway related to fas. This precludes us from using human cell lines for our metastasis experiments, which would require immune-deficient mice such as SCID or nude mice. We selected MXH10/Mo/lpr mice for the present experiments because their characteristics make them better suited for use as a model of metastasis than immune-deficient mice implanted with human cell lines.

Fisher et al.\(^\text{[23]}\) reported that LNs were not an effective barrier against tumor cell progression along the lymphatic system based on the observation that tumor cells were confirmed in the efferent lymphatic vessels after their injection into the lymphatic vessels of the rabbit. This means that tumor cells can flow through the lymphatic system via the marginal sinuses of LNs\(^\text{[17,24]}\), suggesting that lymphatic flow based on the anatomical structure of LNs should be distinguished from the mechanism of tumor metastasis.

Our findings suggest the possibility that tumor cells may undergo hematogenous metastasis at an early stage of LN metastasis, i.e., even before the stage when the infiltration of tumor into the LN is detectable by imaging or aspiration cytology. In other words, false-No metastatic LNs can be a source of hematogenous metastasis. The importance of false-No LNs in distant metastasis is supported by the results of clinical trials suggesting that LN dissection\(^\text{[25,26]}\) and sentinel LN biopsy\(^\text{[27]}\) do not always improve survival in patients with cancer.

The development of new methods to detect and treat metastasis in false-No LNs will be extremely important. In previous studies, we have shown that a lymphatic drug delivery system has great potential in the therapeutic and prophylactic treatment of false-No LNs\(^\text{[25,26]}\), and we expect this technique to be applied in the clinical setting in future.
DECLARATIONS

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Authors’ contributions
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Interpreted the data, reviewed the manuscript: All authors

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