Sympathetic nerve tissue in milky spots of the human greater omentum

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Abstract

Omental milky spots (OMSs), small lymphoid structures positioned in the greater omentum, are involved in peritoneal immune homeostasis and the formation of omental metastases. Sympathetic nerve activity is known to regulate immune function in other lymphoid organs (e.g. spleen and lymph nodes) and to create a favourable microenvironment for various tumour types. However, it is still unknown whether OMSs receive sympathetic innervation. Therefore, the aim of this study was to establish whether OMSs of the adult human greater omentum receive sympathetic innervation. A total of 18 OMSs were isolated from five omenta, which were removed from 3% formaldehyde-perfused cadavers (with a median age of 84 years, ranging from 64 to 94). OMSs were embedded in paraffin, cut and stained with a general (PGP9.5) and sympathetic nerve marker (TH and DBH), and evaluated by bright field microscopy. A T-cell, B-cell, and macrophage staining was performed to confirm OMS identity. In 50% of the studied OMSs, sympathetic nerve fibres were observed at multiple levels of the same OMS. Nerve fibres were represented as dots or elongated structures and often observed in relation to small vessels and occasionally as individual structures residing between lymphoid cells. The current study shows that 50% of the investigated OMSs contain sympathetic nerve fibres. These findings may contribute to our understanding of neural regulation of peritoneal immune response and the involvement of OMSs in omental metastases.

Key words: greater omentum; innervation; metastases; neuroimmunomodulation; omental milky spot; sympathetic.

Introduction

Body cavities are lined by mesothelium and the sub-mesothelial compartment of the pleura, pericardium, and peritoneum (both parietal and visceral) is known to contain lymphoid cell clusters, referred to as milky spots (MSs) (Michailova & Usunoff, 2004). The most prominently present and extensively studied MSs are the ones positioned in the omentum, hence referred to as omental MSs (OMSs). OMSs are cellular aggregates of macrophages, B-cells, and T-cells (Shimotsuma et al. 1991; Liu et al. 2015). They develop during ontogeny (Krist et al. 1997), are found in various animal species including humans (Michailova & Usunoff, 2004), and can be observed in large fixed omental tissue samples after staining with haematoxylin (Schurink et al. 2019) (Fig. 1). OMSs play an important role in the peritoneal immune response: (1) peritoneal fluid is absorbed via openings (stomata) in the omental mesothelial lining and is monitored in OMSs for foreign or pathogenic substances (Van Vugt et al. 1996; Wilkosz et al. 2005; Mepass & Randall, 2017), and (2) OMSs form a gateway for circulating immune cells that are recruited towards the peritoneal cavity during a peritoneal immune challenge (Wijffels et al. 1992). In addition, OMSs are involved in peritoneal metastatic disease; they form primary implantation sites for peritoneal exfoliated cancer cells, which can develop into omental metastases (Krist et al. 1998; Hagiyama et al. 1993; Tsujimoto et al. 1995, 1996).

The autonomic nervous system, and in particular sympathetic nerves, are known to be involved in local regulation of the immune response in lymphoid organs, such as the spleen and lymph nodes (Mignini et al. 2003; Komegae et al. 2018), and in creating a tumour-stimulating microenvironment (Cole et al. 2015). So far only a few studies have...
reported on OMS-associated nerve fibres (Krist et al. 1994; Havrlentova et al. 2017; Yildirim et al. 2010) but none of these have addressed sympathetic fibres specifically. If, like other lymphoid structures, OMSs receive sympathetic innervation, this nerve tissue, or its effectors, could hold potential as therapeutic targets to modify peritoneal immune response or tumor environment. Therefore, the aim of this study is to establish whether OMSs of the adult human greater omentum receive sympathetic innervation.

**Methods**

The greater omenta of five human cadavers were studied. These included three male and two female cadavers with a median age of 84, ranging from 64 to 94 years. Bodies were donated through a body donation programme to the Anatomy Department of the University Medical Center Utrecht, the Netherlands. Informed consent was obtained during life, allowing the use of these bodies for educational and research purposes. No scars were observed in the abdominal region and the available medical records did not list relevant diseases such as cancerous, immune or neurodegenerative disease.

Whole body preservation was accomplished by perfusion with 3% formaldehyde via the femoral artery. The aprons of the omenta were resected from the transverse colon and stored in a 0.1M phosphate buffer, pH 7.4, containing 15% sucrose [phosphate-buffered saline (PBS)/sucrose] at 4 °C until further investigation. One or two large samples of approximately 6 × 4 cm were removed from each omentum. These samples were stained with haematoxylin (Schurink et al. 2019) and evaluated for the presence of OMSs with the aid of a stereomicroscope (Leica EZ4, Nussloch, Germany), using both transmission and incident light. OMSs were identified, removed (with a small margin of surrounding tissue), and placed in PBS/sucrose at 4 °C until further processing for microscopic investigation. As it is known that sympathetic nerve fibres accompany arteries towards their end organs (Mitchel, 1956), omental arterial branches were studied to verify the presence of sympathetic nerve fibres in the selected omental regions. Figure 2 contains schematic images of the selected omental regions, and the amount of isolated OMSs and arterial branches per investigated omentum.

**Immunohistochemistry**

OMSs and omental arteries were processed for paraffin embedding by placing them in increasing percentages of ethanol, xylene, and paraffin. Paraffin blocks were cut on a microtome (Leica 2050 Super Cut, Nussloch, Germany) and 5-μm-thick sections were placed on glass slides, air-dried, and subsequently heat-fixed for 2 h on a slide drying table of 60 °C (Medax, 14801, Kiel, Germany). Each OMS was serially cut and slides were selected at multiple levels for microscopic investigation. Adjacent slides of each OMS level and of each omental artery were stained with antibodies against general [protein gene product 9.5 (PGP9.5)] and sympathetic [tyrosine hydroxylase (TH) and dopamine beta-hydroxylase (DBH)] nerve tissue. If comparable staining patterns were present in TH- and DBH-stained samples, it was assumed that immunoreactive (IR) tissue represented sympathetic nerve tissue. To confirm OMS identity,
additional slides were stained for the presence of B-cells (CD20), T-cells (CD3), and macrophages (CD68), the main cellular constituents of human OMSs (Shimotsuma et al. 1991; Liu et al. 2015). As sympathetic regulation of splenic immune function is known primarily to involve adrenergic activation of T-helper cells (Kin & Sanders, 2006; Olofsson et al. 2012), an additional CD4 antibody staining was performed. All slides were deparaffinized and rehydrated, followed by 20 min of antigen retrieval in 95 °C citrate buffer on a hot plate. After washing in Tris-buffered saline (TBS) with 0.05% Tween (TBS/Tween), sections were pre-incubated for 10 min with 5% normal human or goat serum, with the exception of the ones selected for CD68 staining. After pre-incubation, sections were incubated with primary antibodies in TBS with 3% bovine serum albumin. Table 1 contains details of the primary antibodies. Following incubation, sections were washed with TBS/Tween and incubated for 30 min at room temperature (RT) with undiluted Brightvision Poly-Alkaline phosphatase Goat-anti-Rabbit (ImmunoLogic, Amsterdam, the Netherlands) in the case of PGP9.5, TH, DBH and CD3 staining, or Brightvision Poly-Alkaline phosphatase Goat-anti-Mouse (ImmunoLogic) in the case of CD20, CD68 and CD4 staining. After incubation with secondary antibodies, all sections were washed with TBS and incubated with Liquid Permanent Red (LPR; Dako, Glostrup, Denmark) for 10 min. Subsequently, tissue sections were washed with distilled water and counterstained with haematoxylin, air-dried at 60 °C for 90 min, and coverslipped using Entellan (Merck, Darmstadt, Germany). Human sympathetic trunk sections were included as positive controls for all three nerve markers and human spleen sections for all four immune cell markers. Negative controls were obtained by incubation of sympathetic trunk, spleen, and OMSs slides with TBS-3% bovine serum albumin without primary antibodies. All slides were evaluated using bright field microscopy.

Image acquisition

Brightfield single images were captured at various magnifications using a DM6 microscope with a motorized scanning stage, a DFC7000 T camera, and LASX software (all from Leica).

Results

OMSs could be identified in haematoxylin-stained omental samples of all five cadavers, according to a previous description (Schurink et al. 2019). Of the five studied omenta, 18 OMSs (2–5 per omentum) and 7 omental arterial branches (1–2 per omentum) were sampled and processed for microscopic evaluation. All positive controls

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showed expected IR patterns, whereas all negative controls were clean.

Omental milky spots

All 18 selected OMSs showed specific microscopic morphological OMS features. They were composed of clusters of lymphoid cells, contained blood vessels, and were surrounded by adipose tissue (Shimotsuma et al. 1989; Wilkosz et al. 2005; Liu et al. 2015; Meza-Perez & Randall, 2017).

The medium diameter of these OMSs was 379 µm, ranging from 139 to 1150 µm. TH-IR was present in nine of 18 OMSs (50%). Figure 2 contains schematic drawings of the investigated omenta including the number of studied OMSs and whether they contained sympathetic nerve fibres. Four of five omenta contained TH-IR fibres in 25–100% of their investigated OMSs. Only one cadaver showed no TH-IR fibres in any of its OMSs. TH-IR in OMSs was observed as individual small spots (Fig. 3A–F) or as elongated structures with a dotted appearance resembling varicosities (encircled structures in Fig. 3B). In all TH-IR OMSs, immune labelling was observed at all studied levels of the same OMSs and in comparable regions. TH-IR structures were often closely related to small OMS blood vessels but were also present as individual structures in close proximity to lymphoid cells (Fig. 3). TH-IR patterns were highly comparable with PGP9.5-IR patterns, suggesting that TH-IR represented neural structures. TH-IR patterns were comparable to DBH-IR patterns. Therefore, it was confirmed that these neural structures were adrenergic and not dopaminergic.

Immune cells

All OMSs were composed of B-cells, T-cells, and macrophages, confirming their OMS identity (Fig. 4B–D). CD4 staining was performed on slides of all TH-IR OMSs and CD4-positive cells were clearly present and distributed homogeneously in three of four cadavers with TH-IR OMSs (Fig. 4E). In one cadaver, no CD4-IR cells could be observed in any of the three TH-IR OMSs.

Omental branches

From each of the five cadavers, one or two omental arterial branches were studied, resulting in a total number of seven branches. PGP 9.5-, TH-, and DBH-staining showed comparable patterns and sympathetic nerve fibres were observed in all studied omental arterial branch samples, either as discrete fibres or in bundles. Discrete nerve fibres were present in the adventitia up to the level of the adventitial-medial border in all omental branches, representing intrinsic vessel wall innervation (Fig. 5A,B). In five of seven omental arterial branches, nerve bundles were observed in the adventitia or surrounding connective tissue, representing nerve fibres accompanying these arteries towards more distal omental structures (Fig. 5A,B).

Discussion

This study shows that human omental OMSs contain sympathetic nerve fibres. Often, these nerve fibres were located around small blood vessels and occasionally, individual nerve fibres were observed between lymphoid cells. By means of TH, confirmed by DBH staining, sympathetic fibres were observed in 50% of the studied OMSs. The absence of TH-IR inside the other OMSs is not likely to be explained by either poor or excessive formaldehyde fixation, as TH-IR fibres and bundles have been observed in tissue surrounding these OMSs and omental arterial branches. An age-related decrease of sympathetic tissue in OMS of our relatively old study subjects (median age of 84 years) would be a more plausible explanation, as sympathetic nerve
tissue of secondary lymphoid structures is known to decline with age (Bellinger et al. 1992).

So far, only three other studies have described nerve tissue associated with OMSs. In two of these studies a general nerve marker (S-100) was used and the type of nerve tissue was not further specified (Havlentova et al. 2017; Yildirim et al. 2010). Although these studies visualized small S-100-IR nerve bundles closely related to blood vessels, in between adipocytes or in the surrounding connective tissue, they do not describe small neural structures in close proximity to lymphoid cells. This in contrast to the results of the current study, in which fibres were observed in relation with...
lymphocytes. The third study used transmission electron microscopy in combination with a dopamine (DA) marker to study innervation of OMSs in two patients under the age of 28 years with unaffected omenta. The authors observed non-myelinated nerve fibres in the OMSs of both subjects (Krist et al. 1994). These nerve fibres contained DA-IR vesicles and the authors concluded that the fibres were dopaminergic. However, DA is a precursor for NA and the nerve tissue could represent both dopaminergic and noradrenergic fibres. In the current study the commonly accepted adrenergic nerve marker TH and a marker for dopamine beta-hydroxylase (DBH), the enzyme involved in conversion of DA to NA, were used on adjacent slides of both OMSs and omental arterial branches. TH staining patterns were comparable to DBH staining patterns, suggesting the present nerve tissue to be adrenergic. However, as no double staining was performed, the presence of dopaminergic fibres could not be excluded. Furthermore, the transmission electron microscopic study of Krist et al. (1994) revealed non-myelinated nerve endings to be in close proximity to immune cells (Krist et al. 1994), a finding which is in line with the results of the current study. Previously, this finding led to the suggestion that these nerves might modulate immune cell function by the release of their neurotransmitter. The suggestion that this nerve tissue could modify immune cell function is highly interesting because it opens new opportunities for anti-inflammatory therapies. The concept of neural structures modifying immune function is known as neuroimmunomodulation. It is acknowledged that the sympathetic nervous system plays an important role in regulation of the systemic inflammatory response via the spleen (Komegae et al. 2018). Activation of sympathetic nerve tissue of the splenic plexus results in a cascade of intrasplenic events, starting with the release of norepinephrine (Kees et al. 2003), followed by adrenergic receptor activation on CD4-positive T-cells (Rosas-Ballina et al. 2011; Vida et al. 2011, 2017), which results in production and secretion of acetylcholine and ultimately inhibits the release of pro-inflammatory cytokines from activated macrophages (Borovikova et al. 2000; de

Fig. 4 Presence of immune cells in an omental milky spot. The panel is composed of images of the same omental milky spot (OMS) stained for various immune cells. (a) Overview image of omental tissue containing an OMS (cell-dense region in boxed area) with surrounding adipose tissue. The OMS is positioned directly underneath the mesothelial lining (green dotted line) of the omentum. The abdominal cavity is marked with a large black asterisk. Additional slides were stained to visualize (b) T-cells (CD3), (c) B-cells (CD20), (d) macrophages (CD68), and (e) T-helper cells (CD4). IR structures are stained bright pink and cell nuclei are stained dark blue. Small vascular structures can be discriminated inside the OMS by the presence of erythrocytes which give the vessel lumina a brownish colour (as shown with small black asterisks in d).
Jonge et al. 2005; Kox et al. 2009; Lu & Kwan, 2014). This results in dampening of the systemic inflammatory response. As OMSs are known to contain macrophages (Shimotsuma et al. 1991; Liu et al. 2015) and the current study shows that OMSs receive sympathetic innervation and contain CD4-positive T-cells, all required elements for neuroimmunomodulation appear to be present. Therefore, stimulation of sympathetic fibres of the greater omentum could result in dampening of the peritoneal immune response, which could be beneficial during peritoneal sepsis.

In addition to its possible role in neuroimmunomodulation, sympathetic innervation of OMSs might also be involved in the formation of peritoneal metastases. According to the seed and soil theory, as first proposed by Paget (1889), tumour cells (seeds) have a specific affinity for certain organs or structures (soil); metastasis would only develop if both seed and soil are compatible (Paget, 1889). MSs provide such a favourable soil for certain abdominal tumours (Hagiwara et al. 1993; Tsujimoto et al. 1995,1996; Krist et al. 1998). Why OMSs provide a hotspot for peritoneal metastases has not been fully elucidated yet. Previous studies have suggested a contributory role of OMS-specific mesothelium and blood vessels (Gerber et al. 2006) or adipocytes (Ladanyi et al. 2018). We propose an additional role for sympathetic innervation of OMSs. The sympathetic nerve system and specifically its neurotransmitter noradrenaline are known (1) to contribute to a favourable tumour microenvironment, for example by affecting stromal cells, endothelial cells, and tumour-associated macrophages (Magnon et al. 2013), and (2) to stimulate tumour cell proliferation, via beta (β) adrenergic receptors on tumour cells (Coelho et al. 2017). As since all of these processes are mediated via adrenergic receptors, the effects of β adrenergic blocking as a potential anti-
tumour therapy have been investigated in in vitro, animal, and retrospective patient studies with promising results (Magnon et al. 2013; Coelho et al. 2017; Zahalka et al. 2017). Although further studies are needed to elucidate the exact anti-tumour mechanism, these findings suggest a potential role for β-blockers in the treatment of cancer. As a potential therapeutic application, β-blockers could be administered to the peritoneal cavity, discretely or as an adjuvant in hyperthermic intraperitoneal chemotherapy, which might limit or prevent intraperitoneal spread and growth of tumour cells.

To elucidate the role of sympathetic innervation of OMSs in the peritoneal immune response and metastatic disease, and its therapeutic potential, further studies are needed. These studies should include omenta from younger individuals to avoid age-related decrease of innervation and investigate the amount and location of sympathetic nerve tissue. Furthermore, these studies should investigate the (synaptic) relation of sympathetic nerve fibres to various OMS cell types such as immune and stromal cells, as well as the presence and type of adrenergic receptors on these cells. Although the current study focused on MSs in the greater omentum, similar structures, often referred to as extraomental MSs, can be found in the peritoneum, mesenterium, mediastinum, pericardium, and pleura (Michailova & Usunoff, 2004; Cruz-Migoni & Caamano, 2016). As these structures are involved in local immune activity (Cruz-Migoni & Caamano, 2016), including them in future studies is recommended.

In conclusion, this study showed that human OMSs are sympathetically innervated. Nerve fibres were observed in relation to both blood vessels and lymphocytes. The presence of sympathetic fibres in OMSs contributes to our understanding of how OMS immune function might be regulated during an abdominal immune challenge and why...
they form primary implantation sites for peritoneal exfoliated cancer cells.

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Authors’ contributions

C.C. designed the study, interpreted the data, designed the figures, and wrote the manuscript. B.S. designed the study, interpreted the data, designed the figures, and wrote the manuscript. D.H. performed the experiments, and helped with the interpretation of the data and writing the manuscript. R.B. supervised the project and helped writing the manuscript.

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