Telocytes in pleura: two- and three-dimensional imaging by transmission electron microscopy

Mihail E. Hinescu · Mihaela Gherghiceanu · Laura Suciu · Laurentiu M. Popescu

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Abstract Information about the ultrastructure of connective (interstitial) cells supporting the pleural mesothelium is scarce. Our aim has been to examine whether telocytes (TCs) are present in pleura, as in epicardium and mesentery. TCs are a distinct type of cell, characterized by specific prolongations named telopodes (Tp). We have used transmission electron microscopy (TEM) and electron tomography (ET) to determine whether ultrastructural diagnostic criteria accepted for TCs are fulfilled by any of the cell subpopulations existing in the sub-mesothelial layer in mouse and human pleura. TCs have been identified with TEM by their characteristic prolongations. Tp appear long and moniliform, because of the alternation of podomes (thin segments of less than 0.2 μm) and podoms (small dilations accommodating caveolae, mitochondria, and endoplasmic reticulum). Tp ramifications follow a dichotomic pattern and establish specialized cell-to-cell junctional complexes. TCs, via their Tp, seem to form an interstitial network beneath the mesothelium, covering about two-thirds of the abluminal mesothelial layer. ET has revealed complex junctional structures and tight junctions connecting pleural TCs, and small vesicles at this level in Tp. Thus, pleural TCs share significant similarities with TCs described in other serosae. Whether TCs are a (major) player in mesothelial-cell-induced tissue repair remains to be established. Nevertheless, the extremely long thin Tp and complex junctional structures that they form and the release of vesicles (or exosomes) indicate the participation of TCs in long-distance homo- or hetero-cellular communication.

Keywords Telocytes · Telopodes · Pleura · Electron tomography · Tight junctions · Mouse (C57BL/6) · Human

Introduction

We previously described a new type of cell, which became known as the “interstitial Cajal-like cell” (acronym ICLC), based on the finding that these cells seemed, at first glance, similar to the well-known interstitial cell of Cajal (ICC) found in the gastro-intestinal tract. We described ICLCs in various cavitary (fallopian tube: Popescu et al. 2005a; uterus: Ciontea et al. 2005; Popescu et al. 2006c; gallbladder: Hinescu et al. 2007; heart: Hinescu and Popescu 2005; Hinescu et al. 2006; Popescu et al. 2006a; Mandache et al. 2007; Gherghiceanu et al. 2008; Kostin and Popescu 2009) and non-cavitary (pancreas: Popescu et al. 2005c; mammary gland: Gherghiceanu and Popescu 2005; placenta: Suciu et al. 2007, 2010) organs. However, little by little, it became clear that the ultrastructure of ICLC was (completely) different from that of ICC, and that the difference between these cells was not only semantic, as...
they have different ultrastructure and immunophenotype and therefore are probably functionally distinct (Popescu and Faussone-Pellegrini 2010; free access data is available on www.telocytes.com). Hence, we coined the terms telocyte (TC) for these cells and telopodes (Tp) for their extremely long but thin prolongations in order to prevent further confusion with other interstitial (stromal) cells (e.g., fibroblasts/fibrocytes, mesenchymal cells, or myofibroblasts).

The concept of TCs was rapidly adopted by other laboratories (Bani et al. 2010; Cantarero et al. 2010; Faussone-Pellegrini and Bani 2010; Kostin 2010; Zhou et al. 2010).

Little information exists concerning the ultrastructure of the sub-epithelial tissue of serosae (Carter et al. 2007), including pleura (Hammar 2008). Despite their simple fine morphology, serosae share not only a common embryological origin from splanchnopleural mesenchyme (Lie-Venema et al. 2007; Di Meglio et al. 2009; Gittenberger-de Groot et al. 2010), but also similar functions, based on cytoarchitectural similarities. However, the precise details of interstitial cells composing serosae seem incompletely explored, since a new type of cell has been indentified in omentum (Sakurai et al. 2001), mesentery (Hinescu et al. 2008), and epicardium (Suciu et al. 2009; Gherghiceanu and Popescu 2009, 2010; Popescu et al. 2009, 2010a, 2010b).

Here, we present evidence that ultrastructural criteria (Popescu and Faussone-Pellegrini 2010) usually applied to diagnose TCs are fulfilled by a sub-population of cells present in mouse and human pleura, in the thin submesothelial layer. Despite technology having matured to the point at which the application of electron tomography (ET) specimens in plastic sections has been achievable for almost a decade (McEwen and Marko 2001), little information is available about the cells in tissue (Barcena and Koster 2009). We have used ET in this study in order to examine the particular ultrastructure of TCs with emphasis on Tp connections.

Materials and methods

Mouse parietal pleura samples were obtained from four 8-month-old C57BL/6 mice. Human parietal pleura samples

Fig. 1 Transmission electron microscopy (TEM) images of two serial sections of mouse parietal pleura to illustrate the long telopodes (Tp) and their sinuous trajectory. a Tp (Tp1: 20 μm; Tp2: 24 μm; Tp3: 20 μm) have a convoluted contour suggesting their capacity to compensate for changes in length attributable to pleural expansion (MC mesothelial cells). b TEM image of the adjacent section showing details of the Tp from the boxed area in a. Tp have thin segments (podomeres) alternating with dilatations (podoms), which accommodate mitochondria (m), endoplasmic reticulum (er), and caveolae (arrowheads). Note the shed vesicles (asterisks) in close proximity to Tp. c TEM image of the telocyte network (TC1: 14 μm long, TC2: 15 μm long) beneath the mesothelial cell layer (MC). The boxed area was selected for electron tomography (see Fig. 5). Bars 5 μm (a), 2 μm (b, c)
were obtained from two patients undergoing surgery for non-neoplastic lung diseases. This study was performed with institutional ethical committee approval.

Transmission electron microscopy Transmission electron microscopy (TEM) was performed on small fragments from parietal pleura processed according to a routine Epon-embedding procedure, as previously described (Mandache et al. 2007; Hinescu et al. 2008). Thin sections (about 60 nm) were examined with a Morgagni 286 transmission microscope (FEI Company, Eindhoven, The Netherlands) at 60 kV. Digital electron micrographs were recorded with a MegaView III charge-coupled device (CCD) by using iTEM-SIS software (Olympus, Soft Imaging System, Münster, Germany). Some TEM images were digitally colored by Adobe Photoshop CS3 in order to highlight TCs.

Electron microscope tomography ET was performed by using a Tecnai G2 Spirit BioTwin transmission electron microscope with single-tilt specimen holder (FEI Company) at 100 kV. Electron tomographic data sets were recorded with a MegaView G2 CCD camera (Olympus) in ET mode on 300-nm-thick sections of Epon-embedded mouse parietal pleura. Tomographs were acquired at 1 degree angular increments from $-65^\circ$ to $+65^\circ$ about an axis perpendicular to the optical axis of the microscope, at a
Fig. 3  

a, b TEM images of telocytes (TC) and their overlapping telopodes (Tp) that reinforce the mesothelial cells (MC) at the level of folds formed by mouse parietal pleura. c–e Higher magnifications of boxed areas in a, b. c Telopodes (Tp) in close vicinity of nerve ending (n) beneath mesothelial cells (MC). d, e Telopodes (Tp1-3 in d, Tp1, Tp2 in e) can be seen overlapping each other in the mesothelial folds. Bars 5 μm (a), 2 μm (b), 1 μm (c–e)
magnification of 36,000×. After data alignment, the data sets were reconstructed into a three-dimensional (3D) volume (data collection, reconstruction, and visualization) by using Xplore3D Tomography Suite software (FEI Company). Amira 5.0.1 software (Visage Imaging, Berlin, Germany) was used for 3D imaging.

**Results**

**Electron microscopy**

Electron microscopy of 60-nm thin sections revealed that the main cellular component of interstitial space beneath the mesothelial layer of mouse parietal pleura consisted of numerous TCs (Figs. 1, 2, 3) and few mononuclear cells (Fig. 2a), macrophages (Fig. 2b), and nerve endings (Fig. 3) in a slightly collagenic extracellular matrix. In human parietal pleura, the sub-mesothelial space contained numerous TCs (Fig. 4), mononuclear cells, macrophages, mast cells, and nerve endings in a matrix composed of layers of collagen and elastic fibers (Fig. 4). TCs seemed to form an interstitial network bordering about two-thirds of the abluminal mesothelial layer (Figs. 1, 2, 3, 4), especially at the level of small mesothelial folds (Figs. 1a, 3).

Pleural TCs showed typical features: a small oval-shaped cell body surrounded by a ring of cytoplasm (Figs. 1c, 2a, inset, 3b, 4a, b) and 2–4 extremely long (15–30 μm) but thin prolongations named telopodes (Tp; Figs. 1, 2, 3, 4). The moniliform aspect of Tp resulted from the alternation of two types of segments: podomeres (thin segments of ∼50 nm and therefore below the resolving power of light microscopy) and podoms (dilations of ∼150–300 nm) accommodating caveolae, 1–2 mitochondria, and elements of smooth and/or rough endoplasmic reticulum (Figs. 1b, 2c). Functionally, the trio caveolae/mitochondria/endoplasmic reticulum corresponded to “Ca^{2+}”-uptake/release units” (Popescu et al. 2006b; Gherghiceanu and Popescu 2007).

Tp showed characteristic dichotomous branching pattern (Fig. 3d) with a sinuous trajectory (Fig. 1a, b) that allowed the accommodation of long Tp in a small space (e.g., 20–25 μm long Tp compressed in less than 2 μm). In order to evaluate the extension potential of the Tp, a convolution index was
expressed as their actual length divided by the end-to-end distance in a straight line (Hinescu et al. 2007); a mean value of 7.5 was obtained.

TCs were in close vicinity to nerve endings (Fig. 3c) and established contacts with other interstitial immunoreactive cells (Figs. 2a, b, 4c) or other TCs (Figs. 1c, 3d, e, 4c) in the sub-mesothelial space. TEM showed that TCs formed a network, and that their intricate Tp were often connected through a “plug and socket” complex (Figs. 1c, 3e, 5a).

TCs showed spot-like electron-dense structures on the cortical space of the Tp involved in the “plug and socket” connections and few tight junctions (Fig. 5a). To analyze these junctional complexes, we performed ET.

Electron tomography

ET (0.5 μm³ volume) of the “plug and socket” assembly connecting TCs (Figs. 5b-e, S1, S2) revealed the presence of composite junctions between Tp. The junctional complex was composed of junctional plaques (four clear junctions of about 0.5 μm long/0.2 μm wide are presented in Fig. 5e-e) alternating with free segments (about 0.5 μm long). The junctional plaques showed anfractuous volumes (Figs. 5d, e, S1). The distances between cellular membranes of Tp were variable, with “fusion” points alternating with about 20-nm-wide gaps between membranes at the level of the junctional plaques (Figs. 6, S2). Small vesicles (30–40 nm)
were present within the Tp placed in the center of “plug and socket” assembly (Figs. 5, 6).

Discussion

This paper establishes that telocytes are present not only in mesentery (Hinescu et al. 2008) and in epicardium (Suciu et al. 2009; Gherghiceanu and Popescu 2009, 2010; Popescu et al. 2009, 2010a), but also in mouse and human pleura, suggesting that TCs are a common cellular component of the serosae. From a teleological point of view, the presence of numerous fibroblasts in normal pleura (Carter et al. 2007) is unsuitable because the main function of fibroblasts is to produce collagen fibers, which are by definition non-extensible.

TEM has shown that pleural TCs lie close to nerve endings and immunoreactive interstitial cells (mononuclear cells, macrophages, mast cells). No apparent classical junctions have been observed in between these heterotypic cells, but rather stromal synapses (Popescu et al. 2005b). Of particular interest is that small shed vesicles or exosomes can often be observed in the intercellular space at the level of the stromal synapse suggesting that heterocellular communication might occur (Mathivanan et al. 2010). All these findings suggest that shed vesicles are involved in stromal synapses of TC heterocellular contacts with other interstitial cells, and that the tight junctions support TC homocellular connections.

The present study has revealed that pleural TCs are connected in a network through tight junctions situated at the level of “plug and socket” assembled telopodes (Figs. 5, 6). This elaborate connection between TCs, as revealed by ET, supports a previous hypothesis about long-distance intercellular signaling (Popescu and Faussone-Pellegrini 2010; Popescu et al. 2010b). Data presented in our ET study not only have revealed how difficult the mapping of the spatial distribution of Tp could be by using classical electron microscopy, but have also illustrated the complex interaction between Tp. ET has made possible the examination of TC cellular ultrastructure, opening up perspectives for scrutinizing the 3D architecture of extremely thin and convoluted Tp, which are suspected of playing a key role in cellular function.

In addition to the classical barrier function, the tight junction is emerging as a regulator of cell growth and differentiation and acts as a scaffolding platform for cell signaling and as a docking station for transport vesicles (Schneeberger and Lynch 2004). Moreover, recent studies have shown prominent clusters of proteins involved in vesicular trafficking and synaptic transmission functions at the tight junction domain (Tang 2006; Steed et al. 2010).

A recent study (Gherghiceanu and Popescu 2009) has suggested that TCs are involved in mesothelial renewal. TCs might guide the migration of mesenchymal cells into the mesothelial layer of the epicardium. We have not found such images in this study, but similar mesenchymal cells do indeed exist in close vicinity to TCs beneath the mesothelial layer (Fig. 2a). In view of this, the proposed ideas regarding the origin of regenerated mesothelial cells covering the surface of body cavities, the differentiation of mesothelial cells from macrophages, the proliferation of residual meso-
thelial cells, or the origin of mesothelial cells from fibroblasts (Amari et al. 2002) should be re-examined, because TCs might be important players in mesothelial-cell-induced mesothelial repair. Moreover, the results presented here open doors for theories and new perspective concerning intercellular communication on the basis that “special” junctions have previously been described between mesenchymal cells (Wuchter et al. 2007; Franke et al. 2009).

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