Original Article

Imaging of lysophosphatidylcholine in an induced pluripotent stem cell-derived endothelial cell network

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Introduction: Vascular endothelial cell disorders are closely related to cardiovascular disease (CVD) and pulmonary diseases. Abnormal lipid metabolism in the endothelium leads to changes in cell signalling, and the expression of genes related to immunity and inflammation. It is therefore important to investigate the pathophysiology of vascular endothelial disorders in terms of lipid metabolism, using a disease model of endothelium.

Methods: Human induced pluripotent stem cell-derived endothelial cells (iECs) were cultured on a matrigel to form an iEC network. Lipids in the iEC network were investigated by matrix-assisted laser desorption/ionization (MALDI) time-of-flight (TOF) imaging mass spectrometry (IMS) analysis. Ion fragments obtained by mass spectrometry were analysed using an infusion method, involving precursor ion scanning with fragment ion.

Results: The MALDI TOF IMS analysis revealed co-localized intensity of peaks at \( m/z \) 592.1 and 593.1 in the iEC network. Tandem mass spectrometry (MS/MS) analysis by MALDI-imaging, in conjunction with precursor ion scanning using an infusion method with lipid extracts, identified that these precursor ions were lysophosphatidylcholine (LPC) (22:5) and its isotype.

Conclusion: The MALDI-imaging analysis showed that LPC (22:5) was abundant in an iEC network. As an in vitro test model for disease and potential therapy, present analysis methods using MALDI-imaging combined with, for example, mesenchymal stem cells (MSC) to a disease derived iEC network may be useful in revealing the changes in the amount and distribution of lipids under various stimuli.

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1. Introduction

Lipids are involved in various cellular functions and play a role in homeostasis and disease states [1–3]. Several lipid species in lipid classes such as fatty acids (FAs), phospholipids (PLs), and ceramides (CMs) have been proposed as useful biomarkers in early identification of disease [4,5]. Endothelial cells (ECs) are recognized as multifunctional cells involved in: regulating the transfer of oxygen; nutrients; coagulation; vascular tone; angiogenesis; stimulating smooth muscle cells and fibroblasts, and the vasculature composition of normal and/or tumor microenvironments [6–8]. External stresses, such as oxidative, glycative or lipoxidative, can alter lipid profiles in ECs compared to the homeostatic state [6,9]. Lipidomics has become a field of research as a result of technological advances in analytical tools of mass spectrometry (MS), nuclear magnetic resonance (NMR), high-performance of liquid chromatography (HPLC), and imaging mass spectrometry (IMS) [10,11]. These techniques identify unique lipid species that indicate potential adverse cardiovascular events. Specifically,
phosphatidylcholine (PC) (32:0) and sphingomyelin (SM) (16:0) have been associated with mortality in cardiovascular disease (CVD) [12], and lysophosphatidylcholine (LPC) (18:1) and (18:2) has been shown to be associated with incidents of CVD [13]. The dysfunction of ECs can result in atherosclerosis [14] and IMS has revealed increased levels of LPC, phosphatidic acid (PA), PC and SM in aortic tissue [15,16].

Induced pluripotent stem cell (iPSc)-derived differentiated cells have been used for drug discovery, drug repositioning and toxicological screening of drugs [17]. Currently, iPSc-derived cell therapy is being used to investigate age-related macular degeneration (AMD), Parkinson’s disease (PD), fibro dysplasia ossificans progressiva (FOP) and spinal cord injury (SCI) [18–21]. In rats, disease models of idiopathic or heritable pulmonary hypertension (IPAH/HPAH), X-linked adrenoleukodystrophy (X-ALD), lipopolysaccharide-induced acute lung injury using induced pluripotent stem cell-derived endothelial cells (iEcs) have been reported [22–24]. The investigation of IPAH/HPAH in this model revealed a reduction in the expression of several genes that may play a role in the effectiveness of treatment by affecting drug repositioning [22].

The first report investigating lipid accumulation was conducted in a blood brain barrier (BBB) model consisting of human brain microvascular iEcs derived from childhood cerebral-ALD (cc-ALD) Ecs. This study showed increased lipid accumulation as evaluated by Oil-Red-O staining in the BBB [23]. The specific lipid classes or clinical meanings of an increase in lipids in iEcs remains, however, unknown.

Lipids are closely related to clinical information in disease states, therefore disease models using iEcs may be useful in investigating the role of lipids in disease states to provide patients with more effective therapeutic treatments.

In the present study, an iEcs network culture model was generated from human normal iEcs. The two-dimensional lipid distribution was examined in the iEcs network by IMS. A co-culture model with lung tissue from human pulmonary mesenchymal stem cells (HPMSC) and the iEcs network was also analyzed. The findings from this present study indicate that IMS analysis of lipids in a disease-derived iEcs network model may be beneficial in identifying patients at risk of CVD and providing more effective treatment for these patients.

2. Materials and methods

2.1. Cell culture of iPSc-derived endothelial cell network on indium/ tin oxide-coated (ITO) glass slides

Human iPSc derived endothelial cells (iEcs) were obtained from Cellular Dynamics Inc (iCell®, Wisconsin, U.S.A). The iEcs were cultured in medium of Vasculi® VEGF Comp Kit (Vifone Cell Technology, CA, U.S.A) as previously [25]. For IMS analysis, iEcs were cultured in Matrigel® (Corning, NY, U.S.A) on indium/tin oxide-coated (ITO) glass slides (Sigma–Aldrich, U.S.A) in Quadruperc® 4-Compartment Culture Dish (Greiner Bio One, Austria). Briefly, the culture dish was kept at 4 °C, and the ITO glass slide was placed in the culture dish. The matrigel was then placed onto the cold ITO glass, and extended thinly by spinning on ice. The ITO glass slide was then incubated in CO2 at 37 °C for 30 min. Following this, 80% confluent cultured iEcs were washed three times with phosphate buffered saline (PBS) at room temperature (RT), then treated with 0.025% phenol free trypsin (NAKARAI, Japan) for 2–3 min, and harvested by cell scraper. The collected iEcs were centrifuged at 1000 rpm for 5 min, and stored in 1.5 × 10^5/ml iEcs were placed onto matrigel-mounted ITO glass slides, and incubated in CO2 at 37 °C for 1 h. After 1 h incubation, 6 ml of 37 °C culture medium was added to the culture dish to cover the surface of cells on the ITO glass slide. After 48 h culture (CO2 at 37 °C), the ITO glass slides were removed from the culture dish and washed three times by cold saline (Otsuka Pharmaceutical, Japan). The iEcs network was identified by microscopic examination (BZ-X, KEYENCE, Japan) and the glass slide snap frozen at −80 °C with desiccant of silica until use. Control samples of matrigel on ITO glass slides were prepared in a similar manner, minus iEcs. The HPMSCs were cultured in mesenchymal stem cell medium (MSCM) in accordance with the manufacturer’s instructions (ScienCells Research laboratory, CA, U.S.A). For coculture with the iEcs network and HPMSC (ScienCells Research laboratory, CA, U.S.A), the iEcs network formation was prepared prior to adding HPMSCs. The HPMSCs were labelled with CellLight™—mitochondria GFP, BacMam 2.0 (Fischer Scientific, U.S.A) for 45 min according to the manufacturer’s instructions, and treated with 0.125% trypsin—EDTA (Invitrogen), and harvested. Following centrifugation at 1000 rpm for 5 min, HPMSCs were transfected into the iEcs network on the ITO glass slide at the concentration of 1 × 10^5 with 50%/50% volume culture medium of Vasculi®/MSCM and incubated for 48 h. After 48 h of culture, the ITO glass slides were removed from the culture dish and washed three times with cold saline. The contact areas between the iEcs network and green fluorescent protein (GFP)-HPMSCs were identified using a fluorescence microscope (BZ-X, KEYENCE).

2.2. MALDI TOF IMS

The MALDI TOF IMS analysis was performed with an iMScope-TRIO® (SHIMADZU, Japan) equipped with a 355 nm Nd:YAG laser, as previously described [26]. The ITO glass slide stored at −80 °C was left at RT to air dry. The iEcs network was traced on the screen monitor using a microscope equipped with iMScopeTRIO® (SHIMADZU, Japan). The inside of the traced area was then analyzed with MALDI-imaging analysis. The ITO glass slide was then homogenously coated with a matrix (2-hydroxy-5-methoxybenzoic acid: DHB) using an iMLayer® (SHIMADZU, Japan). The MS data were acquired in positive ion mode in the mass range m/z 500 to 1000. A laser beam diameter was set at 5 μm, and mass resolving power was 10,000 at m/z 1000 used in MS and tandem MS/MS scans. Principal component analysis (PCA) and hierarchical clustering analysis (HCA) of image patterns was performed as previously described [26]. The target m/z was determined after comparison with the m/z peaks derived from the control matrigel without iEcs. A sample ITO glass slide without iEcs but coated with matrigel was prepared in a similar manner and used as a control.

2.3. Extraction of lipids in Ecs network for LC-MS and database research

The ITO glass slide was washed three times in cold saline, and the matrigel, including the iEcs network, was treated with cell recovery solution according to the manufacturer’s instructions (Corning NY, U.S.A.). The iEcs were then collected. After sonication of the cells, lipids were extracted by the Bligh-Dyer method as previously described [26]. Samples were stored at −80 °C after sealing under nitrogen. To identify the unknown lipids co-localized between the iEcs network and green fluorescent protein (GFP)-HPMSCs by MALDI-imaging. The lipidView™ database with QTRAP5500 mass spectrometer (Sciex Inc, Framingham, MA USA) was used to determine specific ions.
2.4. Statistical analysis

The PCA was performed for the cumulative contribution ratio of PC > 80% in each image and the contribution of PC value in each group > mean PC value of each image. The HCA was performed in accordance with the Euclidean distance and the Ward Method. The PCA, HCA, and generation of dendrograms were performed using iMScopeTRIO® (SHIMADZU, Japan) software.

3. Results

3.1. MALDI-imaging analysis of iEC network

Formation of the iEC network or matrigel without iEC on an ITO glass slide was confirmed by microscopic examination before being stored at −80 °C (Fig. 1a–c). The ITO glass slides were air dried, and the iEC network traced using microscopy equipped with iMScopeTRIO® (SHIMADZU, Japan). The inside of the traced area was analyzed in regards to the iEC network (Fig. 1d) or in the matrigel without the iEC network (Fig. 1f). The MS analysis of the iEC network showed various intensities in m/z peaks including those derived from the matrigel and from the iEC network (Fig. 1a). High intensity of the peaks at m/z 545.0 and 585.0 were determined from DHB (Fig. 2b). The PCA analysis based on two-dimensional image patterns identified 24 groups (Supplemental data 1). PCAs including peaks at m/z 592.1, 593.1, 632.3, and 728.1 were co-localized to the iEC network (Fig. 3a). The PCA analysis showed 13 clusters (Supplemental data 2). The following HCA measurements were co-localized with the iEC network: HCA10 of m/z 592.1 and 593.1; HCA12 of m/z 632.3, and HCA13 of m/z 672.4. Among the distribution of m/z 592.1, some spots were high in the iEC network as shown yellow to red dots (Fig. 3b). Furthermore, dendrogram of the HCA clusters demonstrated that these m/z peaks formed close trees (Fig. 4). These peaks of m/z were not observed on matrigel without the iEC network.

3.2. Tandem MS/MS analysis of peaks of m/z co-localized to iEC network

To confirm the lipids co-localized with the iEC network, tandem MS/MS analysis was performed on peaks of m/z 592.1, 593.1, 632.3, 672.4 and 728.1 by MALDI-imaging analysis. From the tandem MS/MS analysis on m/z peaks of 592.1 and 593.1 as precursor, a peak at m/z 184.1, a specific fragment of phosphoryl choline ion, was obtained (Fig. 5a and b). From other precursor m/z peaks at 632.3, 672.4 and 728.1, valuable fragments m/z could not be obtained (Supplemental data 3). To identify target lipids of m/z peaks at 592.1 and 593.1, lipid extract from the iEC network underwent analysis using the infusion method. Phosphoryl choline m/z 184-containing phospholipids were determined via a database search of LipidView™ equipped in QTRAP5500. Peaks at m/z 592.1 and 593.1 were identified as LPC 22:5 (569.34) + Na (23) and the isotype. Other peaks at m/z 632.3, 672.4, and 728.1 could not be identified in the database. Tandem MS/MS analysis for the peak of ion precursor at m/z 728.1 showed a fragment at m/z 669.4, with the difference between 728.1 and 669.4 being 58.7. The neutral loss (NL) of 58.7 Da could be NL of 59 Da, and NL of 59 Da is a well known fragment from the phosphatidylcholine (PC) species of trimethylamine [N(CH3)3]. Therefore, m/z 728.14 might be PCs, however, no lipids matched the database.

3.3. Co-culture model of the iEC network and HPMSC

After forming the iEC network, GFP-HPMSCs were added with culture medium. The GFP-HPMSCs were observed in the iEC network line or joint. A representative pattern of GFP-HPMSCs in the iEC network is shown (Supplemental data 4). MALDI-imaging analysis were done on co-culture model. Intensity of the peaks of m/z 592 and 672 co-localized to iEC network were identified but it was difficult to distinguish the metabolic differences between GFP-HPMSC and iEC, or a GFP-HPMSC effect on iEC network (Supplemental data 5 and 6).

4. Discussion

The MALDI-imaging analysis of the iEC network demonstrated differences in the relative intensity reflecting various peaks of m/z. Tandem MS/MS analysis by MALDI-imaging of the iEC network and infusion method analysis using lipid extract identified LPC (22:5), and it was abundant in iEC network. A proposed model of co-culture with an iEC network and HPMSCs was also demonstrated.

Fig. 1. Cell culture and preparation of slides for MALDI-imaging. Matrigel was mounted onto a sterilized ITO-coated glass slide and placed in a 4-compartment culture dish. Human iPSC-derived vascular endothelial cells (iECS) were cultured in matrigel for 2 days to form an iEC network (a). An ITO glass slide mounted with matrigel but without iECs underwent culture medium. The GFP-HPMSCs were observed in the iEC network (b). The iEC network (e) or the matrigel without the iEC network (f) were again confirmed by microscopic examination before being stored at −80 °C with desiccant. On the day of MALDI-imaging analysis, the slide was air dried at RT (d). The iEC network (e) or the matrigel without the iEC (f) were then stored at −80 °C. After forming the iEC network, GFP-HPMSCs were added with culture medium. The GFP-HPMSCs were observed in the iEC network line or joint. A representative pattern of GFP-HPMSCs in the iEC network is shown (Supplemental data 4). MALDI-imaging analysis were done on co-culture model. Intensity of the peaks of m/z 592 and 672 co-localized to iEC network were identified but it was difficult to distinguish the metabolic differences between GFP-HPMSC and iEC, or a GFP-HPMSC effect on iEC network (Supplemental data 5 and 6).
One of the lysophospholipids generated from glycerophospholipid metabolism cleaved by phospholipase A2 or A1 is LPC [28,29]. It is well known that LPC plays an important role in cell signaling, migration, proliferation and the release of cytokines in various types of cells including: ECs; adipocytes; hepatocytes; immune cells, and neuronal cells [30]. It has been shown that ECs can be activated by the following: LPC; induced monocyte chemoattractant protein-1 (MCP-1); interleukin-8 (IL-8); regulated on activation, normal T cell expressed and secreted (RANTES) expression, cytotoxicity or apoptosis, and oxidative stress [31–33].
Interestingly, LPC has a potent property that can result in trans differentiation of human aortic ECs into innate immune cells via up-regulation of cytosolic calcium and mitochondrial reactive oxygen species (ROS) [34]. Furthermore, it has been reported that LPC may play a role in the induction of histone acetylation levels at H3K14 in 19 trained immunity genes in ECs. The acetylation status of H3K14 is currently considered a biomarker or therapeutic target of CVD, further supporting the need to consider LPC as a biomarker for CVD [35].

As a biomarker, LPC has been observed in the plasma, serum and other fluids in the following diseases: bronchoalveolar lavage fluid (BALF); lipid extraction of tissue in cystic fibrosis; idiopathic pulmonary fibrosis (IPF); chronic obstructive pulmonary disease (COPD) CVD; hepatic cancer; lung cancer patients, and asthma.
In a murine bleomycin-induced pulmonary fibrosis model, LPC was reported to be associated with the pathophysiology of this disease [30,43]. In the present study, LPC (22:5) was identified in an iEC network. Several LPCs, including LPC (14:0), (16:0), (18:0), (18:2), (20:4), (22:3) and (22:6) have been reported [30], however, few reports have identified LPC (22:5) [29,40,44]. In patients with hepatocellular carcinoma (HCC) LPC (22:5) levels were decreased which may indicate poorly differentiated HCC [40]. In a murine lung fibrosis experimental model, an increase in LPC (22:5) was observed in the lung, however, there was no increase in systemic LPC [29]. In addition, LPC (22:5) was produced from phospholipase A activity of PCs, and also by autotaxin (ATX) dependently, which was also elevated in the lung in the murine lung fibrosis experimental model [29]. The relationship between ECs and LPC is complex. The stimulation by the combination of LPC and ATX treatment has been shown to increase the permeability of human umbilical vein EC (HUVEC), and immunohistochemistry (IHC) has revealed that the area of vascular leakage was co-localized with ATX protein expression in a murine model of renal fibrosis [44]. The findings from the present study show that LPC (22:5) was abundant in ECs, and that injury to ECs may cause increased levels of LPC at the site of injury. This increase in LPC levels may exacerbate the vascular metabolic state.

The MALDI-imaging analysis can obtain two-dimensional data. Studies have been conducted investigating LPC distribution in injured ischemic rat brain, ischemic porcine kidney and restenosis in injured rat carotid artery [45–47]. These studies reported an increase in LPC (18:1) at the injured site, and that the LPC levels were increased compared to the control by the time course (day 0 to day 14 of injury) [47]. In addition, LPC (20:4) and (22:6) have also been identified in murine normal airway by nanospray desorption electrospray ionization MS coupled to shear force microscopy [48], however, there have been no reports of LPC (22:5) by MS imaging. The present study focused on the expression of LPC in an iEC network, composed of an iEC network developed on matrigel by cell culture. The difference between the present study and previous reports are that previous studies focused on tissue sections including various cells, and on larger cell areas other than the iEC network, and different matrix or ion-modes were used for analysis of LPC expression.

The present study reports findings of LPC expression using a co-culture cell model with HPMSCs and an iEC network. Several studies have reported metabolic changes of stem cell transplant by IMS analysis [49,50]. IMS analysis has revealed that glial stem cell transplants to the murine brain lowers PA, diacylglycerol (DAG) metabolism [49], and bone marrow derived mesenchymal stem cells (BM-MSC) in injured left soleus muscle of rat altered protein expression in a murine model of renal injury. This increase in LPC levels may exacerbate the vascular permeability of human umbilical vein EC (HUVEC), and non-targeted lipidomics. Biochem Biophys Res Commun 2018;504:576–81.

None of the authors have any conflict of interests.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.reth.2020.03.007.

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5. Conclusion

The findings of the present study indicate that a model of co-culture iEC network and HPMSC may be beneficial in determining the expression of lipids in a disease state. Furthermore, LPC (22:5) was abundant in the iEC network, and MALDI-imaging analysis was useful in identifying the distribution of lipids in the iEC network.
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