Protective Effect of Se-Methylselenocysteine on Elaidic Acid-Induced Inflammation in Human Arterial Endothelial Cells

Jizhu XIA1, Xiaorong XIA1, Wenyuan WANG1, Jiyi XIA2 and Mingxing LI1

1Department of Ultrasound, The Affiliated Hospital of Southwest Medical University, Luzhou, Sichuan, 646000, China
2Southwest Medical University, Luzhou, Sichuan, 646000, China

(Received June 9, 2020)

Summary This study was designed to investigate the anti-inflammatory effect of Se-methylselenocysteine (MSC) on elaidic acid (9t18:1, EA) induced human arterial endothelial cells (HAECs). MTT and flow cytometry were used to determine cell viability and cell apoptosis respectively. Western blotting was used to assess protein expression of intercellular adhesion molecular 1 (ICAM-1), E-selectin, interleukin-8 (IL-8), endothelial nitric oxide synthase (e-NOS) and phospholipases A2 (PLA2), while enzyme-linked immunosorbent assay (ELISA) was performed to examine the secretion level of nitric oxide (NO). In the cell viability assay, EA significantly decreased cell viability when compared with negative control (NC) group, and MSC effectively reversed this adverse effect, especially at the concentration of 200 μmol/L with 24 h incubation. Also, the same concentration of MSC prevented HAECs cell apoptosis induced by EA. In addition, we found that the expression of ICAM-1, E-selectin, IL-8 and PLA2 were significantly increased and e-NOS decreased in EA group compared with NC group. Inhibition of PLA2 promoted ICAM-1, E-selectin and IL-8 expression in HAECs induced by EA. And MSC down-regulated the secretion of NO level in EA-induced HAECs. Based on these results, we concluded that MSC activated PLA2 which regulated the expression of ICAM-1, E-selectin and IL-8 to protect inflammation induced by EA in HEACs.

Key Words Se-methylselenocysteine, elaidic acid, endothelial cell, inflammation, PLA2

Trans fatty acids (TFAs) have attracted wide attention on cardiovascular health for a long time. Epidemiology evidence have demonstrated that industrial TFAs (I-TFAs) promotion the expression of inflammatory cytokines with the main component of elaidic acid (9t18:1, EA) (1). Also, published clinical studies have demonstrated that I-TFAs are closely related to endothelial cell damage, cardiovascular diseases (CVDs) and insulin resistance (2, 3).

The function of endothelial cells is mainly related to angiogenesis and formation of vascular wall and maintenance of vascular integrity and permeability. Harvey et al. (4) reported that EA induce endothelial cell dysfunction by elevating levels of inflammation cytokines such as intercellular adhesion molecular 1 (ICAM-1). Iwata et al. (5) found that EA promote endothelial cell injury by increasing nuclear factor-κB (NF-κB) expression and influencing the production of insulin-mediated nitric oxide (NO) level. Furthermore, I-TFAs also increasing the risk of atherosclerosis by inhibiting the synthesis of polyunsaturated fatty acids (PUFA) in arterial cells, and EA can incorporate into phospholipids on endothelial cell membranes and then trigger the inflammation response (6).

Cell membrane plays a crucial role in cell biological activities and energy metabolism. Phospholipids, the most important part of cell membrane, are involved in lots of cell activities, such as information transmission, external material substance and exchange. Studies demonstrated that phospholipids could be degraded into arachidonic acid (AA) and other lysophospholipids by stimulation of various factors. These degradation products will finally contribute for vascular injury (7).

Phospholipase is an important substance on cell membrane that regulates phospholipid synthesis and metabolism. Phospholipases A2 (PLA2) is one of the major isomers of phospholipases which are divided into three types, secretory phospholipases A2 (sPLA2), cytoplasmic phospholipases A2 (cPLA2) and calcium-independent phospholipases A2 (iPLA2). It has been reported that PLA2 could hydrolyze phospholipids, generate AA and other free fatty acids (7).

Selenium, one of the most essential non-metal trace elements in human, closely related to many diseases related to chronic inflammation, like inflammatory bowel diseases, rheumatoid arthritis, and atherosclerosis (8, 9). Previous studies have proved that selenium may block many key processes to protect vascular injury. The underlying mechanism of selenium on protect endothelial cell inflammation could be summarize as reduction of recruitment of inflammatory cells by inhibition activation of p38 MAPK (10), and inhibition of

E-mail: xiajizhu2@yeah.net
endothelial dysfunction and endothelial cell apoptosis by increasing antioxidase activity (11), reverse NO release (12), down-regulating caspase-3 and Bax expression (13). But few articles investigated the role of selenium on activity of phospholipases and the following anti-inflammation process.

Therefore, the present study was designed to explore the effect of Se-methylselenocysteine (MSC) on bio-activity of PLA2, and to evaluate following influence to human arterial endothelial cells (HAECs) inflammation induced by EA.

**MATERIALS AND METHODS**

**Regents.** HAEC cells were bought from Cell Bank of the Chinese Academy of Science (Shanghai, China). Elaidic acid, MTT, DMSO, Se-methylselenocysteine (MSC; purity ≥95%) were obtained from Sigma (St. Louis, USA). Dulbecco’s modied eagle medium (DMEM), fetal bovine serum (FBS), total RNA extraction kit (TRizol), annexin V-FITC apoptosis detection kit and enhanced chemiluminescence (ECL) kit were bought from Invitrogen (Carlsbad, USA). Trans Script assay kit, real time PCR assay kit, polyclonal ICAM-1, E-selectin, IL-8, e-NOS, iPLA2, sPLA2, cPLA2 and p-cPLA2 antibodies were obtained from Santa Cruz Biotechnology (California, USA).

**Cell culture and treatment.** Cells were maintained in an incubator at 37°C with 5% CO2, supplemented with DMEM medium containing 10% fetal bovine serum, 100 μg/mL penicillin G, and 100 μg/mL streptomycin sulfate, and medium was changed every 3 d.

**MTT assay.** MTT assay was curried out to assess cell viability. Briefly, cells were divided into 5 groups: negative control (NC) group, elaidic acid (EA) group (50 μmol/L), EA+MSC group (50 μmol/L+100 μmol/L), 50 μmol/L+200 μmol/L, 50 μmol/L+400 μmol/L). The concentrations of EA and MSC used in this study were according to previous study (14, 15). Three thousand cells per well were seeded into 96-well plates and incubated for 24 h and 48 h with different reagents. After incubating with MTT for 4 h, dimethyl sulfoxide was added in the dark and co-culture with cells for another 10 min. Finally, OD value was recorded at 570 nm.

**Flow cytometry.** Cells were cultured with treated with different reagents as the same as MTT assay. Forty-eight hours later, flow cytometric assay was used to determine apoptosis through an Annexin V-FITC apoptosis detection kit following the manufacturer’s protocol.

**Western blot.** Cells were divided into following groups: NC group, EA group; EA+MSC group (50 μmol/L+100 μmol/L), 50 μmol/L+200 μmol/L, 50 μmol/L+400 μmol/L); EA+MSC+IB group: cells were pretreated with different inhibitors for 1 h, and then cultured with EA and MSC for 24 h. IB refers to inhibitors for PLA2, including PYR (cPLA2 inhibitor), DTT (sPLA2 inhibitor) and BEL (iPLA2 inhibitor). The incubation time and concentration of inhibitors were followed by the previous report (16).

Proteins were extracted using RIPA buffer containing cocktail protease inhibitor. Total concentration of protein was measured by BCA assay. Protein samples (20 μg) were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) gel and blotted onto ployvinylidine diufoe (PVDF) membranes. After blocked with 5% non-fat milk for 1 h, membranes were subsequently incubated with primary antibodies and HRP-conjugated secondary antibody. Signals were measured by the enhanced chemiluminescence (ECL) kit.

**Enzyme-linked immunoassay (ELISA).** Effect of EA and MSC on NO and prostaglansin E2 (PGE2) release level in HAECs were determined by ELISA assay. HAECs were seeded into 96-well plates at the density of 1×10⁵ cells per mL. Cells were incubated with EA and MSC as the same as described in MTT assay, and the culture supernatants was prepared for ELISA assay according to the manufacturer’s protocol using NO and PGE2 ELISA kits.

**Statistical analysis.** All assays were performed in triplicate except for the MTT (n=6). All experimental data were presented as mean±standard deviation. One-way ANOVA followed by Tukey’s test was used to determine differences between groups with SPSS 22.0. p-values of less than 0.05 were considered statistically significant.

**RESULTS**

**Cell viability**

Cell viability of HAECs was detected by MTT assay after cells were treated with different agents for 24 h or 48 h. As the results showed in Fig. 1, the cell viability decreased signiicantly after treated with 50 μmol/L EA compared with NC group (p<0.05), and the cell viability were 68.21±2.55% and 57.85±6% at 24 h and 48 h respectively. Compared with the EA group (48 h), the cell viability was dramatically increased when treated with EA+100 μmol/L MSC for 48 h (57.28±3.17% vs. 69.58±1.91%, p<0.05). Moreover, the result showed that comparison with other treated group: EA+
200 μmol/L MSC for 24 h resulted the highest viability (82.67 ± 2.28%) of HAECs. However, with the concentration of MSC increased to 400 μmol/L, the cell viability decreased significantly at 24 h (62.85 ± 1.84%) or 48 h (51.66 ± 2.27%) when compared with the EA group (68.34 ± 2.51% and 57.55 ± 3.17% at 24 h and 48 h respectively) (p < 0.05).

Cell apoptosis

After incubated with EA for 24 h, the apoptosis rate of HAECs was significantly increased when compared with NC group (23.47 ± 1.47% vs. 8.72 ± 1.47%, p < 0.05). After stimulated with EA + MSC (100 μmol/L and 200 μmol/L), the apoptosis rate was decreased dramatically in a dose-dependent manner (15.47 ± 3.48% and 11.32 ± 1.09% respectively) compared with EA group (p < 0.05). However, there was no significant change of the apoptosis rate between EA + MSC 400 μmol/L group and EA group (p > 0.05) (Fig. 2).

Expression of inflammatory cytokines

The expression level of inflammatory related factors, ICAM-1, E-selectin, and IL-8 were significantly increased (p < 0.05), and the expression level of e-NOS was decreased in the EA group when compared with the NC group (p < 0.05). Meanwhile, MSC significantly reversed the adverse effect of EA in EA + MSC group. And pretreatment cells with PLA2 inhibitors for 1 h, and then incubated with EA + MSC, the expression level of ICAM-1, E-selectin (except EA + MSC + PYR group) and IL-8 significantly increased when compared with EA + MSC group (p < 0.05). Also, the expression level of e-NOS decreased significantly in the EA + MSC + BEL group compared with EA + MSC group (p < 0.05) (Fig. 3).

Expression of PLA2

As shown in Fig. 4, the expression of all four kinds of PLA2 (iPLA2, sPLA2, cPLA2 and cPLA2 phosphorylation (p-cPLA2)) were dramatically increased in HAECs after incubated with EA for 24 h compared with NC group (p < 0.05), and obvious up-regulation of sPLA2 was found in EA group. Moreover, the expression of PLA2 significantly decreased after treatment of EA + MSC (100 μmol/L and 200 μmol/L), especially for the expression level of p-cPLA2 at the group treated with...
Secretion of NO and PGE2

Figure 5 represent the effect of EA and MSC on the secretion level of NO in HAECs. EA significantly decreased the secretion of NO compared with NC group \( (p<0.05) \). MSC (100, 200 and 400 \( \mu \)mol/L) significantly reversed the effect of EA. The EA+MSC (200 \( \mu \)mol/L) group shows strongest effect among all EA+MSC groups \( (p<0.05) \). Figure 6 shows PGE2 levels in HAECs. Compared with NC group, EA significantly increased the secretion of PGE2 \( (p<0.05) \). After treatment with MSC, PGE2 section decreased significantly compared with EA group \( (p<0.05) \).

DISCUSSION

In the present study, we found that EA significantly decrease the viability of HAECs and increase apoptosis, and MSC dose-dependently reverse this adverse effect. The most appropriate concentration and duration of MSC to incubate with HAECs might be 200 \( \mu \)mol/L and 24 h. The apoptosis rate of HAECs increased obviously when co-incubated with EA+400 \( \mu \)mol/L. MSC may due to cytotoxicity of MSC at high concentration \( (17) \). Reported studies approved that the most two common existence forms of selenium, inorganic selenium (such as sodium selenite) and organic selenium (such as methylseleninic acid and Se-methylselenocysteine) may exert bioactive or cytotoxic effects in vitro strictly depend on compound, concentration and model used \( (14, 18) \). We use MSC in our model because increasing studies supported that MSC has stronger bioactivity than other forms of selenium in vitro \( (19) \).

Atherosclerosis is one of the leading causes of death over the world, and the dysfunction of endothelial cells was considered as the initial factor of atherosclerosis. Selenium has been proved as a crucial non-metal trace element in the process of metabolism of life. Although
Se-methylselenocysteine Prevent HAECs Inflammation

limited numbers of prospective research reported inconsistent result on the protect effect of selenium on atherosclerosis, the benefit of selenium and selenoproteins supplementation has been verified to protect experimental vascular injury (20). In the present study, we stimulated HAECs with EA, one of the major TFA forms in human diet, and then investigate the protect effect of MSC on HAECs inflammation. Our results revealed that MSC could reverse the pro-inflammatory effect of EA. In addition, the secretion level of PGE2 (a down-stream product of AA) in EA group was significantly higher than EA + SMC groups (Fig. 6). This effect of MSC may due to its anti-inflammation power through AA. The oxidation products of AA, including hydroperoxycisataetraenoic acids (HpETEs) could promote inflammation, and reports found that HpETEs can be reduced by glutathione peroxidase 4 (GPx4) (21), one of the family members of GPxs, which containing about 4 g of selenium atoms per 1 mole (17). Furthermore, GPx4 also benefit for anti-inflammation by inhibiting the nuclear factor kappa-B (NF-kB) pathway, reducing the expression of lipoxigenase and cyclooxygenase (22–25). Despite the abundant anti-inflammatory mechanisms of MSC, but whether and how MSC inhibit the generation of AA still unclear.

PLA2 is widely distributed in the biological cell membrane. The main function of PLA2 is hydrolyzes phospholipids and then generate free fatty acids, including AA and lysolceithin, which induce endothelial dysfunction (7). Previous study reported that cPLA2 could be activated by stimulating of H_{2}O_{2} via ERK pathway (26), and it could also be activated by IL-6, TNF-α and histamine (27, 28). Also, many studies found stronger activity of cPLA2 in inflammatory cells than non-inflammatory cells (29). Ma et al. (30) reported that up-regulation of sPLA2 involved in inflammatory pathological injury to hepatopancreas caused by phoxim. Sato et al. (31) revealed that enhanced decomposition of cardiolipin following mitochondrial iPLA2 activation might mediate trogitazone-induced mitochondrial membrane permeability transition.

In the present study, we found that EA induced the expression of PLA2, and EA + MSC groups were negatively correlated with the PLA2 expression level. The expression of ICAM-1, E-selectin, and IL-8 were also significantly decreased in EA + MSC groups compared with EA group (p < 0.05). However, groups pretreated with PLA2 inhibitor (DTT, BEL and PYR) were positively correlated with the expression of those inflammatory cytokines (ICAM-1, IL-8) compared with the EA + MSC group (p < 0.05). Meanwhile, the E-selectin level in groups pretreated with inhibitor (DTT and BEL) were higher than EA + MSC group (p < 0.05), and e-NOS level in group pretreated with inhibitor (DTT and PYR) showed no significant difference with EA + MSC group (p > 0.05), and inhibitor BEL depress expression of e-NOS compared with EA + MSC group (p < 0.05). Based on these results, we assume that PLA2 may involved in the ICAM-1, IL-8 and E-selectin expression in EA induced HAECs inflammation. This results were coordinate with Hu’s report (32), and we first revealed that PLA2 may not participate in EA induced e-NOS expression in HAECs.

Nitric oxide, known as the main chemical substance for maintaining normal vascular homeostasis, was produced by e-NOS. Previous studies have shown that selenium supplementation for 3 d enhanced NO release in acetylcholine-induced rats (33). Lu et al. also reported that selenium supplementation increased NO secretion in homocysteine-induced endothelial injury (34). In our study, we also find that NO and e-NOS were significantly decreased after treated with EA, and MSC significantly induced NO release and e-NOS expression compared with EA group. Interestingly, our results demonstrated that only PLA2 inhibitors of BEL suppressed expression of e-NOS compared with inhibitor free group. These results suggested that PLA2 pathway may not involve in secretion of NO in our model, and further potential underlying mechanisms need to be clarified.

Nonetheless, there are some limitations in this in vitro study. The antioxidant system in which selenium involved is very complex, other solutes except PLA2 may interact the biological system. So, more in vivo and clinical trials should be carried out to better verify theprotective effect of SMC on EA induced HAECs.

Authorship
Jizhu Xia designed this study. Xiaorong Xia and Wenyuan Wang collected and analyzed the data. Jiyi Xia and Mingxing Li wrote and approved the final version of the manuscript. All authors read and approved the manuscript for publication.

Disclosure of state of COI
The authors declare that they have no conflict of interests.

Acknowledgments
We thank the National Nature Science Foundation of China (No. 81501481); Key technologies R&D program of Sichuan Province and Luzhou City (14JC0130).

REFERENCES
1) Mozaffarian D, Clarke R. 2009. Quantitative effects on cardiovascular risk factors and coronary heart disease risk of replacing partially hydrogenated vegetable oils with other fats and oils. Eur J Clin Nutr 63: S22–S33.
2) Wang Y, Jacome-Sosa MM, Proctor SD. 2012. The role of ruminant trans fat as a potential nutraceutical in the prevention of cardiovascular disease. Food Res Int 46: 460–468.
3) Willett WC, Stampfer MJ, Manson JE, Colditz GA, Speizer FE, Rosner BA, Sampson LA, Hennekens CH. 1993. Intake of trans-fatty-acids and risk of coronary heart disease among women. Lancet 341: 581–585.
4) Harvey KA, Walker CL, Pavlina TM, Xu ZD, Zaloga GP, Siddiqui RA. 2008. Trans-fatty acids induce pro-inflammatory responses and endothelial cell dysfunction. Br J Nutr 99: 723–731.
...