HMG-CoA reductase inhibitors induce apoptosis of lymphoma cells by promoting ROS generation and regulating Akt, Erk and p38 signals via suppression of mevalonate pathway

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Statins, the inhibitors of 3-hydroxy-3-methylglutaryl coenzyme A (HMG-CoA) reductase, are widely used cholesterol-lowering drugs. Convincing evidence indicates that statins stimulate apoptotic cell death in several types of proliferating tumor cells in a cholesterol-lowering-independent manner. The objective here was to elucidate the molecular mechanism by which statins induce lymphoma cells death. Statins (atorvastatin, fluvastatin and simvastatin) treatment enhanced the DNA fragmentation and the activation of proapoptotic members such as caspase-3, PARP and Bax, but suppressed the activation of anti-apoptotic molecule Bcl-2 in lymphoma cells including A20 and EL4 cells, which was accompanied by inhibition of cell survival. Both increase in levels of reactive oxygen species (ROS) and activation of p38 MAPK and decrease in mitochondrial membrane potential and activation of Akt and Erk pathways were observed in statin-treated lymphoma cells. Statin-induced cytotoxic effects, DNA fragmentation and changes of activation of caspase-3, Akt, Erk and p38 were blocked by antioxidant (N-acetylcysteine) and metabolic products of the HMG-CoA reductase reaction, such as mevalonate, farnesyl pyrophosphate (FPP) and geranylgeranyl pyrophosphate (GGPP). These results suggests that HMG-CoA reductase inhibitors induce lymphoma cells apoptosis by increasing intracellular ROS generation and p38 activation and suppressing activation of Akt and Erk pathways, through inhibition of metabolic products of the HMG-CoA reductase reaction including mevalonate, FPP and GGPP.

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It is known that malignant lymphoma (ML), a heterogeneous disease with highly variable clinical course and prognosis, is the most prevalent type of adult leukemia.1–3 Most patients with MLs in clinical course are aggressive and soon after diagnosis require intensive treatment.4,5 Both the defective balance between pro- and anti-apoptotic molecules, and aberrant upregulation of prosurvival mechanism have been shown to be related to resistance of ML cells to radiation therapy and chemotherapy.6,7 Previous clinical studies have demonstrated that symptomatic ML can be effectively treated with purine analogs, glucocorticoids, alkylating agents or monoclonal antibodies. However, some patients with relapsed or refractory disease have limited therapeutic options. Therefore, there is an urgent need to discover less toxic and more effective drugs for ML patients.

Inhibitors of 3-hydroxy-3-methyl glutaryl coenzyme A (HMG-CoA) reductase (statins) are used to treat hypercholesterolemia. Convincing evidence from both in vitro and in vivo data has demonstrated that statins exert pleiotropic actions beyond their lipid-lowering effects, including immune regulation8 and cancer prevention.9,10 Statins have been demonstrated to induce cell cycle arrest and cell death in various cancer cells such as multiple myeloma cells,11 pancreatic cancer cells,12 non-small lung cancer cells,13 waldenstrom macroglobulinemia cells,14 glioblastoma cell lines15 and HT29 cells.16 A recent study has shown that simvastatin inhibits proliferation of MCF-7 cells in parallel with an increase in reactive oxygen species (ROS) production.17 Another lipophilic statin, atorvastatin, has also been shown to elevate levels of myocardial protein oxidation and lipid

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Abbreviations: ROS, reactive oxygen species; ΔΨm, mitochondrial membrane potential; HMG-CoA, 3-hydroxy-3-methylglutaryl coenzyme A; Mev, mevalonate; FPP, farnesyl pyrophosphate; GGPP, geranylgeranyl pyrophosphate; DCFH-DA, 2’,7’-dichlorodihydrofluorescein diacetate; HO, Hoechst 33342; PI, propidium iodide; DAPI, 4,6-diamidino-2-phenylindole; NAC, N-acetylcysteine; JC-1, 5,5’,6,6’-tetrachloro-1,1’,3,3’-tetraethylbenzimidazolylcarbocyanine iodide; SOD, superoxide dismutase; CAT, catalase

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peroxidation. Moreover, a high-dose of atorvastatin induces oxidative DNA damage in human peripheral blood lymphocytes.

Previous studies have demonstrated that cancer cells produce higher levels of ROS than normal cells and this contributes to cancer progression. To maintain ROS at tolerable physiological levels, cancer cells possess an antioxidant defense system that includes glutathione and glutathione-dependent enzymes such as superoxide dismutase and catalase to eliminate ROS. Increased ROS generation selectively sensitizes oncogenically transformed and cancer cells, but not non-transformed cells, to cell death, indicating that neoplastic cells are more vulnerable to increased intracellular oxidative stress. Given these previous findings, we hypothesized that statins exert at least some of their cytotoxic effects by increasing oxidative stress depending on cell type.

In the present study, we investigated the effects of statins including atorvastatin, fluvastatin and simvastatin on survival of lymphoma cells such as A20 and EL4 cells, and explored the potential underlying mechanism. We demonstrated that statins induce lymphoma cells apoptosis by increasing intracellular ROS generation and p38 activation and suppressing activation of Akt and Erk pathways, through inhibition of metabolic products of the HMG-CoA reductase reaction including mevalonate, farnesyl pyrophosphate (FPP) and geranylgeranyl pyrophosphate (GGPP).

Results

Fluvastatin-induced cytotoxicity in lymphoma cells. The effects of statins on viability of peripheral blood mononuclear cells (PBMCs) and lymphoma cell lines (A20 and EL4 cells) were determined using the EZ-CyTox Cell Viability Assay Kit as described in method section. Cells were incubated with atorvastatin, fluvastatin or simvastatin at concentrations ranging from 0–5 μM for 24 and/or 48 h, respectively. Our results revealed that, statins at low concentration of 1.25 and 2.5 μM exerted minimal effects on the ability of primarily isolated PBMCs after treatment for 24 h, even they significantly inhibited the cell viability at 5 μM. However, each statin significantly decreased the viabilities of A20 and EL4 cells after treatment of 24 h, even at lowest concentration of 1.25 μM. Furthermore, statins inhibited viability of lymphoma cells in a dose- and time-dependent manner. However, fluvastatin showed higher cytotoxicity towards lymphoma cells than atorvastatin or simvastatin. Even at 24 h, fluvastatin (5 μM) inhibited the viability of A20 cells and EL4 cells by ~50% and 40%, respectively (Figure 1a). Therefore, fluvastatin was selected to use throughout the following experiments.

After treatment with fluvastatin (0–20 μM) for 24 h, cell death was then examined by using trypan blue staining. As shown in Figure 1b, fluvastatin markedly induced cell death of A20 cells and EL4 cells in a dose-dependent manner. Even at 2.5 μM, fluvastatin induced ~25% of cell death of two cancer cells.

Apoptosis was involved in fluvastatin-induced cytotoxicity towards lymphoma cells. To explore apoptosis whether involved in fluvastatin-induced cell death in lymphoma cells, we next investigated the amount of sub-G1 DNA in cancer cells that treated with fluvastatin using flow cytometry. As shown in Figure 2, the treatment of lymphoma cells with fluvastatin resulted in the increased accumulation of cells in the sub-G1 phase in a dose-dependent manner. To further elucidate apoptosis stage of cancer cells induced by fluvastatin, Hoechst 33342 (HO)/propidium iodide (PI) double staining method was used. The plasma membrane of viable cells is only slightly permeable to HO, leading to light-blue nuclear fluorescence. However, HO efficiently crosses the plasma membrane of apoptotic cells due to increased membrane permeability, causing bright-blue fluorescence of the nuclei. On the other hand, PI only penetrates cells with damaged membranes, leading to bright-red fluorescence of nuclei. Thus, intact light-blue nuclei (HO+/PI−), condensed/fragmented bright-blue nuclei (HP+/PI−), condensed/fragmented pink nuclei (HP+/PI+), intact pink nuclei (HP+/PI+) were considered to indicate viable, early apoptotic, late apoptotic (secondary necrotic) and necrotic cells, respectively. Following treatment with fluvastatin at concentrations of 5 and 10 μM for 24 h, HO/PI double staining showed mainly necrotic cell death of primary PBMCs. However, massive apoptotic, but not necrotic, cell death was observed in both A20 and EL4 cells (Figures 3a-b). To explore the dose-response effects of fluvastatin on apoptosis, both cancer cells were incubated with fluvastatin at concentrations ranging from 0–20 μM for 24 h. Annexin V-FITC/PI staining showed that fluvastatin induced apoptosis in cancer cells in a dose-dependent manner (Figures 3c and d). Taken together, these findings suggest that apoptosis is involved in fluvastatin-induced cytotoxicity in lymphoma cells.

Fluvastatin-induced nuclear condensation. Apoptotic morphological changes were assessed by staining with 4,6-diamidino-2-phenylindole (DAPI) and fluorescence microscopy. After treatment with fluvastatin at concentrations of 5 and 10 μM for 24 h, marked morphological changes induced by fluvastatin in a dose-dependent manner were observed, such as nuclear condensation, nuclear fragmentation, and apoptotic bodies (Figures 4a and b). Transmission electron microscopy was utilized to further assess the features of the apoptotic cell death induced by fluvastatin. The control cells exhibited normal cell morphology, but features of apoptotic cells such as chromatin condensation and apoptotic bodies were observed after treatment with fluvastatin at concentrations of 5 and 10 μM for 24 h (Figure 4c). To explore the effect of fluvastatin on cell apoptosis, apoptosis was also detected by DNA fragmentation assay. DNA fragmentation was significantly increased after treatment with fluvastatin in a dose-dependent manner (Figure 4d). Taken together, these data indicated that nuclear condensation and DNA fragmentation were involved in fluvastatin-induced apoptotic death of A20 and EL4 cells.

Fluvastatin treatment led to decreased mitochondrial membrane potential (ΔΨm). To further document the involvement of mitochondrial dysfunction in lymphoma cell apoptosis induced by fluvastatin, we next measured ΔΨm in A20 cells with flow cytometry analysis and JC-1 staining. Cells were incubated with fluvastatin at concentrations
ranging from 0–20 μM for 12 h and analyzed by using flow cytometry. As shown in Figures 5a and b, with increase in the concentration of fluvastatin, the number of cells emitting green fluorescence (apoptosis) increased from 18.24% in control cells to 54.42% in those treated with fluvastatin at 20 μM. Statistical analysis from three independent data showed that treatment with fluvastatin significantly reduced DC in a dose-dependent manner, as indicated by a decrease in red/green ratio, from 0.3082 ± 0.0031 in the control to 0.0592 ± 0.0022 in fluvastatin-treated cells (Figure 5c).

**Effects of fluvastatin on apoptosis-related molecules.** To further explore the molecular mechanism contributing to statins-induced apoptosis, the expression of apoptosis-related proteins was examined by western blot analysis. As shown in Figure 6a, the expression of cleaved caspase-3 was remarkably enhanced in both A20 and EL4 cells
following treatment with atorvastatin, fluvastatin or simvasta- 
tin at 5 \( \mu \text{M} \) for 12 h, respectively. In addition, fluvastatin 
(5 \( \mu \text{M} \)) greatly increased the expression of cleaved caspase-3 
in both two cancer cells in a time-dependent manner (Figure 6b). We also treated A20 cells with fluvastatin at 
concentrations ranging from 0–10 \( \mu \text{M} \) for 12 h. The expres- 
sions of cleaved caspase-3 and cleaved PARP, the well-
known characteristics of apoptosis, were significantly 
increased in a dose-dependent manner (Figure 6c). The 
apoptosis defects are mainly determined by a defective 
balance among pro- and anti-apoptotic members of the Bcl-2 
family, often related to resistance of cancer cells to 
chemotherapy\(^6\). The expression of Bax, a pro-apoptotic 
protein, was increased while expression of Bcl2, an anti-
apoptotic protein, was decreased in fluvastatin-treated A20 
cells (Figure 6c). Moreover, the activity of caspase-3 in A20 
cells was also observed to increase in a dose-dependent 
manner after treatment with fluvastatin (Figure 6d).

Furthermore, Akt pathway is the major anti-apoptotic 
molecular that confer the survival advantage and resistance 
of cancer cells against various chemotherapeutic agents.\(^{25}\) We first investigated whether fluvastatin (5 \( \mu \text{M} \)) downregulated 
constitutive Akt activation in lymphoma cells. As shown in 
Figure 6e, constitutive phosphorylation (activation) of Akt was 
suppressed by fluvastatin in a time-dependent manner. We 
also analyzed the activation of MAPK cascades including p38 
and Erk in A20 cells. We found that fluvastatin markedly 
increased phosphorylation of p38 MAPK and decreased the 
phosphorylation of Erk pathway in a time-dependent manner, 
respectively (Figure 6e). These results indicate that
fluvastatin can suppress the activation of Akt and Erk pathways, but promote the activation of p38 MAPK pathway in lymphoma cells.

Oxidative stress was involved in fluvastatin-induced cytotoxicity. To investigate the involvement of oxidative stress in fluvastatin cytotoxicity, we examined the important oxidative stress marker, intracellular ROS levels, in lymphoma cells following treatment with fluvastatin at concentrations ranging from 0–20 μM for 6 h. As shown in Figure 7, treatment of lymphoma cells with fluvastatin significantly increased intracellular ROS generation in a dose-dependent manner, suggesting the potential involvement of oxidative stress in the cytotoxic action of fluvastatin. To further explore the signaling mechanism of ROS in fluvastatin-induced cytotoxicity towards lymphoma cells, we incubated A20 cells with fluvastatin (5 μM) in the presence or absence of the thiol antioxidant N-acetylcysteine (NAC). Cells were treated with 5 mM NAC for 2 h before and during 12 h exposure to fluvastatin, cell viability, western blotting and DNA fragmentation were then analyzed. As shown in Figure 8a, NAC could significantly block increase in the expression of cleaved caspase 3 and p38 MAPK regulated by fluvastatin, whereas the fluvastatin-inhibited activation of Akt and Erk pathway.
were markedly blocked by NAC. In addition, both cell viability inhibition and DNA fragmentation induced by fluvastatin were remarkably suppressed by NAC (Figures 8b and c).

**Mevalonate pathway contributes to fluvastatin-induced apoptosis in lymphoma cells.** To examine the signaling mechanism for fluvastatin-induced cytotoxicity towards A20 cells, we incubated cells with fluvastatin in the presence or absence of mevalonate (Mev, 200μM), GGPP ammonium salt (GGPP, 10μM) or FPP ammonium salt (FPP, 10μM). Western blotting data in Figure 8a showed that the increase in expression of cleaved caspase 3 and p38 MAPK regulated by fluvastatin were markedly suppressed, whereas the fluvastatin-inhibited activation of Akt and Erk pathway were markedly blocked by Mev, FPP or GGPP. In addition, both cell viability inhibition and DNA fragmentation induced by fluvastatin were remarkably suppressed by Mev, FPP or GGPP (Figures 8b and c). Taken together, these data indicate that mevalonate pathway may contribute to fluvastatin-induced apoptosis in lymphoma cells.

**Discussion**

Convincing evidence from both in vitro and mouse model data suggest that statins can be used as a potential cancer therapeutic depending on the type of cancer cell, but the
effects of statins on ML cells and related mechanism have been veiled. To clarify this issue, we examined whether different statins (atorvastatin, fluvastatin and simvastatin) induce cytotoxicity in A20 cells and EL4 cells. Our results revealed that statins markedly suppressed the viability of lymphoma cells in a dose- and time-dependent manner. However, fluvastatin showed more cytotoxicity towards lymphoma cells than other two statins, by increasing intracellular ROS generation and p38 activation and suppressing activation of Akt and Erk pathways, through inhibition of metabolic products of the HMG-CoA reductase reaction including mevalonate-FPP and GGPP.

Previous studies have reported that statins can induce cell death in various cancer cells in a cell type-dependent manner.11,13,15,17,26 These previous data are consistent with our results showing that statins, especially fluvastatin, induced significant inhibition of the viability of lymphoma cells. We next documented that apoptosis was responsible for fluvastatin-induced cytotoxicity towards A20 cells using flow cytometry, HO/PI double staining, TEM, DNA fragmentation and annexin V-FITC staining, indicating that fluvastatin treatment directly induced an apoptotic death in lymphoma cells.

The increasing knowledge about the processes that regulate apoptosis has identified several targets, which can be used as specific cell death markers, including the changes in mitochondrial membrane potential.27 The decrease in mitochondrial membrane potential and the release of cytochrome C into the cytoplasm may occur upon apoptotic stimulation. Caspase-9 is activated due to the combination of released cytochrome C and apoptotic protease activating factor-1, thereby processing other caspase members, including caspase-3 and caspase-7, to initiate a caspase cascade, which leads to apoptosis.28,29 Despite we did not directly examined the release of cytochrome C, our results revealed a dose-dependent decrease in the mitochondrial membrane potential and increase in the activation of caspase-3 in A20 cells following treatment with fluvastatin, suggesting that the mitochondrial pathway is also involved in fluvastatin-induced cell apoptosis. Because PARP is one of the main cleavage targets of caspase-3,30 we next examined the cleavage of PARP. As expected, the cleavage of PARP was observed in lymphoma cells, suggesting that cells were undergoing apoptosis.31 On the other hand, the apoptosis defects are mainly determined by a defective balance among pro- and anti-apoptotic members of the Bcl-2 family, often related to resistance of CLL B-cells to chemotherapy.32 In this study, the expression of Bax was increased but that of Bcl-2 was decreased in fluvastatin-treated lymphoma cells, indicating that the resistance of lymphoma cells to apoptosis can be blocked by the addition of fluvastatin.

Several signaling pathways, including Akt, Erk and p38 were showed to be important for cell cycle progression and proliferation.33–35 In the present study, treatment with fluvastatin markedly suppressed the activation of Akt and Erk. However, the phosphorylation of p38 pathway was markedly increased by fluvastatin in A20 cells (Figure 6e), indicating the involvement of these three pathways in fluvastatin-induced apoptotic death in lymphoma cells. The hypothesis was further supported by previous studies. For instance, statin can suppress the activation of Akt, a major prosurvival pathway, in cancer cells.36 Furthermore, p38 pathway-mediated apoptosis was also observed in different cell types.37,38 In addition, Erk activation is essential for

![Figure 5](image-url)
carcinogenesis, and constitutively activated Erk is found in a variety of human cancers. Recent studies indicate that increased intracellular ROS generation may be involved in statin-induced cytotoxicity in MCF-7 breast cancer cells. Furthermore, atorvastatin treatment is associated with elevated levels of myocardial protein oxidation and lipid peroxidation in a mouse model. These previous studies are at least partly consistent with our data showing the potential involvement of intracellular ROS generation in fluvastatin-induced cytotoxicity towards lymphoma cells. Inhibition of HMG-CoA reductase by statins is limiting for the biosynthesis of not only cholesterol, but also other important isoprenoid intermediary metabolites such as dolichols, and the electron transport chain proteins heme A and ubiquinone. A lack of these non-steroid isoprenoids, which are related to antioxidant status, might cause oxidative stress. Moreover, neoplastic cells are more vulnerable to increase in ROS level because they function with a heightened basal level of ROS-mediated signaling.

Combining these previous studies with our observations in this study, we hypothesize that statins, especially fluvastatin, cause a breakdown of the antioxidant defense system and thereby increasing the accumulation of intracellular ROS to levels that exceed the cell’s metabolic capabilities to maintain an acceptable physiological range. In support of this idea, a well-known antioxidant, NAC, suppressed the DNA fragmentation and cytotoxicity induced by fluvastatin (Figure 8). Other studies have also suggested that statins can induce cytotoxicity in an oxidative stress-dependent manner. For example, atorvastatin has been demonstrated to induce oxidative DNA damage in human peripheral blood lymphocytes. Furthermore, increased intracellular ROS production is responsible for lovastatin-induced cell death of k-ras-transformed thyroid cells. Although they use a different experimental system, these studies partially support our results showing that fluvastatin-induced cytotoxicity is accompanied by an increase in intracellular ROS generation in A20 cells. These results further indicate that increased accumulation of...
intracellular superoxide is involved in the death of lymphoma cells induced by fluvastatin.

Statins are known to lower cholesterol by inhibiting HMG-CoA reductase, thereby blocking the mevalonate pathway. Besides reducing cholesterol biosynthesis, inhibition of mevalonate by statins also leads to a reduction in the synthesis of isoprenoids such as FPP and GGPP. However, these intermediates are involved in the positive modulation of several non-steroid isoprenoids (for example, heme A, coenzyme Q10 and dolichols) that are related to antioxidant status, and a reduction in these non-steroid isoprenoids increases intracellular oxidative stress by disrupting the antioxidant defense system in certain transformed and cancer cells, particularly by inhibiting biosynthesis of isoprenoid antioxidants such as CoQ10 and dolichol. This idea is further supported by our results showing that fluvastatin-induced cytotoxicity, apoptotic signaling and DNA fragmentation were significantly decreased by addition of Mev, FPP, GGPP (Figure 8).

In conclusion, the present study demonstrates that treatment with HMG-CoA reductase inhibitors induced apoptotic death in ML cells, by increasing intracellular ROS generation and p38 activation and suppressing activation of Akt and Erk pathways, through inhibition of metabolic products of the HMG-CoA reductase reaction including mevalonate, FPP and GGPP. Moreover, decrease in mitochondrial membrane potential was also contributed in HMG-CoA reductase inhibitor-induced apoptosis. This new understanding supports the development of mevalonate pathway-targeted therapy with HMG-CoA reductase inhibitors as direct therapeutic agents for treating ML.

Statins induce lymphoma cell apoptosis

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Materials and Methods

Reagents. Atorvastatin (calcium salt, (C33H34FN2O5)2Ca 3H2O) was obtained from Pfizer (Groton, CT, USA). Fluvastatin (sodium salt, C24H25FNNaO4) and simvastatin (sodium salt, C24H25O3 Na) were purchased from Calbiochem (La Jolla, CA, USA). DCFH-DA, HO, PI, DAPI, NAC, Mev, GGPP ammonium salt and FPP ammonium salt were purchased from Sigma-Aldrich Co (St. Louis, MO, USA). 5,5,6,6'-tetrachloro-1,1'-3,3'-tetraethyl-benzimidazolyl-carbocyanine iodide (JC-1) was purchased from Molecular Probes Inc. (Eugene, OR, USA). Antibodies against cleaved caspase-3, poly ADP-ribose polymerase (PARP), cleaved PARP, Bax, Bcl2, Akt, phospho-Akt (Ser473), p38 MAPK, phospho-p38 MAPK, Erk1/2, phospho-Erk1/2, β-actin and HRP-conjugated goat anti-rabbit IgG were from Cell Signaling Technology (Beverly, MA, USA). A sensitive western blotting luminal reagent was obtained from Santa Cruz Biotechnology Inc. (Santa Cruz, CA, USA).

Cell culture. A20 and EL4 lymphoma cells (Korean Cell Line Bank, Seoul, Korea) were cultured in RPMI 1640 medium (BioWhittaker Inc., Walkersville, MD, USA) containing 10% fetal bovine serum, 100 U/ml penicillin and 100 µg/ml streptomycin (all from BioWhittaker Inc., Walkersville, MD, USA) at 37 °C in a 5% CO2 incubator. In experiments described below, the medium was exchanged for RPMI 1640 medium containing 2% fetal bovine serum. PBMCs, isolated from C57BL/6 mice, were incubated as described above and used as a normal control.

Cell viability assay. Cell viability was analyzed using EZ-CyTox Cell Viability Assay Kit (Doell Lab Service Co Ltd, Seoul, Korea) according to the manufacturer’s instructions. In brief, dispense 90 µL of cell suspension (1 × 10^4 cells) in a 96-well plate. After 4 h, 10 µl of various concentrations of statins or pharmacological reagents dissolved in medium or medium alone was added into corresponding wells. The plate was incubated for indicated time, 10 µl of EZ-CyTox Cell Viability Assay Kit solution was then added to each well of the plate followed by 2 h-incubation at 37 °C in a 5% CO2 incubator. In experiments described below, the medium was exchanged for RPMI 1640 medium containing 2% fetal bovine serum. PBMCs, isolated from C57BL/6 mice, were incubated as described above and used as a normal control.

DNA flow cytometric analysis. After treatment with fluvastatin, cells were harvested, washed twice with ice-cold phosphate-buffered saline (PBS), fixed with 75% ethanol at –20 °C overnight, washed again and then incubated with RNase
A (25 μg/ml, Bio Basic Inc., Markham, Ontario, Canada) at 37 °C for 30 min. Cells were washed once with PBS and incubated with PI (50 μg/ml) for 30 min at room temperature in the dark. The cells were resuspended in 500 μl PBS and subjected to flow cytometry on a Cytomics FC500 flow cytometer, followed by data analyses using Summit version 5.2 software (Beckman Coulter). The cells with sub-G0/G1 peak were evaluated as DNA degradation caused by apoptosis.

Detection of changes in mitochondrial membrane potential by JC-1. Mitochondrial membrane potential was evaluated by using JC-1 (Molecular Probes) staining and flow cytometry analysis. The JC-1 powder was dissolved in dimethyl sulfoxide to make a stock solution at concentration of 5 mg/ml. Lymphoma cells were incubated with JC-1 (1 μM) at 37 °C for 15 min in the dark, washed and resuspended in 500 μl PBS. Cells were then subjected to flow cytometry on a Cytomics FC500 flow cytometer (Beckman Coulter). Data analyses were performed using Summit version 5.2 software (Beckman Coulter, Fullerton, CA, USA). Data were analyzed using Summit version 5.2 software (Beckman Coulter, Fullerton, CA, USA).

Annexin V and PI staining. Phosphatidylserine on the cell surface was detected with Annexin V-FITC Apoptosis Detection Kit I (BD Pharmingen, San Diego, CA, USA) according to the manufacturer’s instructions. In brief, cells were washed twice with cold PBS and resuspended in 100 μl annexin-V binding buffer, followed by incubation with FITC-conjugated annexin V and PI for 15 min at room temperature in the dark. The cells were resuspended in 500 μl binding buffer and analyzed by Cytoomics FC500 flow cytometer (Beckman Coulter, Fullerton, CA, USA). Data were analyzed using Summit version 5.2 software (Beckman Coulter, Fullerton, CA, USA).

Figure 8 Fluvastatin-induced cytotoxicity was reversed by mevalonate, FPP, GGPP, and NAC. (a) A20 cells were incubated with fluvastatin (5 μM) in the presence or absence of mevalonate (200 μM), FPP (10 μM), GGPP (10 μM) and NAC (5 mM) for 12 h, and the expression of cleaved caspase-3 and phosphorylation of Akt, p38 and Erk were examined by western blotting. (b and c) A20 cells were incubated with fluvastatin (5 μM) in the presence or absence of mevalonate (200 μM), FPP (10 μM), GGPP (10 μM) and NAC (5 mM) for 24 h. (b) The cell viability was quantified by using EZ-CyTox Cell Viability Assay Kit and expressed as the percentage of the viability of resting cells. Results are presented as mean ± S.E.M. of three separate experiments conducted in duplicate. **P<0.01 versus cells treated with fluvastatin. (c) The DNA fragmentation was examined by using DNA fragmentation assay. Lane 1, Marker; Lane 2, fluvastatin; Lane 3, fluvastatin + mevalonate; Lane 4, fluvastatin + FPP; Lane 5, fluvastatin + GGPP; Lane 6, fluvastatin + NAC

DNA fragmentation assay. After treatment, cells were harvested in a 1.5 ml Eppendorf tube, washed with PBS and resuspended in 400 μl lysis buffer (10 mM Tris pH 8.0, 10 mM NaCl, 10 mM EDTA, 1% SDS and 100 μg/ml Proteinase K) and incubated at 65 °C overnight. Seventy-five microlitres of potassium acetate (8 M) was then added and the samples were incubated at 4 °C for 15 min. Seven hundred and fifty microlitres of chloroform was added into the Eppendorf tube, which was then mixed vigorously and centrifuged (12000 × g) at room temperature for 10 min. The supernatant was transferred into a new Eppendorf tube and 750 μl ethanol was added, followed by overnight incubation of the sample at -20 °C. DNA was acquired by centrifugation (12000 × g, 10 min) of the sample, washed, dried, and dissolved in 50 μl TE buffer (10 mM Tris-HE, 1 mM EDTA, pH 8.0). Five micrograms of DNA were analyzed on 2.0% agarose gel.

**HO/PI double staining.** Cells were incubated with either 1 μg/ml HO or 5 μg/ml PI at 37 °C, 5% CO2 for 15 min in the dark and collected by centrifugation. The cell pellets were fixed in 4% formaldehyde, washed with ice-cold PBS, resuspended, and a fraction of the suspension was smeared on a slide. The slide was air dried, mounted with VECTAShield mounting medium and examined under a DMI 4000 fluorescence microscope (Leica, Wetzlar, Germany). Morphological assessment of apoptosis and necrosis was performed as follows: intact light-blue nuclei (HO+/PI–), condensed/fragmented bright-blue nuclei (HP++/PI–), condensed/fragmented pink nuclei (HO+/PI–), intact pink nuclei (HO+/PI+), and condensed/fragmented pink nuclei (HO+/PI–). FPP (10 μM) was then added and the samples were incubated at 4 °C for 15 min. Seven hundred and fifty microlitres of chloroform was added into the Eppendorf tube, which was then mixed vigorously and centrifuged (12000 × g) at room temperature for 10 min. The supernatant was transferred into a new Eppendorf tube and 750 μl ethanol was added, followed by overnight incubation of the sample at -20 °C. DNA was acquired by centrifugation (12000 × g, 10 min) of the sample, washed, dried, and dissolved in 50 μl TE buffer (10 mM Tris-HE, 1 mM EDTA, pH 8.0). Five micrograms of DNA were analyzed on 2.0% agarose gel. The DNA fragmentation was examined by using DNA fragmentation assay. Lane 1, Marker; Lane 2, fluvastatin; Lane 3, fluvastatin + mevalonate; Lane 4, fluvastatin + FPP; Lane 5, fluvastatin + GGPP; Lane 6, fluvastatin + NAC

Cell viability (%) of A20 cells was quantified by using EZ-CyTox Cell Viability Assay Kit and expressed as the percentage of the viability of resting cells. Results are presented as mean ± S.E.M. of three separate experiments conducted in duplicate. **P<0.01 versus cells treated with fluvastatin.
Western blotting assay. Cells were washed twice with ice-cold PBS and resuspended in 150 μl RIPA cell lysis buffer (Sigma-Aldrich Co, St Louis, MO, USA), mixed completely, incubated on ice for 30 min, and followed by centrifugation 12 000 × g for 20 min at 4 °C. Supernatants were then stored at – 80 °C until use. Protein concentrations were determined using the Bio-Rad Protein Assay (Bio-Rad Laboratories, Hercules, CA, USA). Equal proteins (30 μg lane) were separated on SDS-polyacrylamide gel electrophoresis and transferred onto polyvinylidene difluoride membrane (Immobilon-P; Millipore, Bedford, MA, USA). Membranes were then blocked with 5% nonfat milk, washed briefly, incubated with primary antibodies at 4 °C overnight, and then incubated with corresponding HRP-conjugated secondary antibodies for 1 h at room temperature. Protein bands were visualized by incubating membranes with chemiluminescence reagents before exposure to X-ray film.

Caspase-3 activity assay. Cells were harvested by centrifugation and washed twice with PBS. The cell pellets were lysed with 100 μl lysis buffer (BD Pharmingen, Cat. No. 559759) and the lysates were incubated with Ac-DEVD-pNA (fluorometric caspase-3). After incubation for 2 h at 37 °C, the fluorescence was measured with a Fluoroskan Asp (Thermo Fisher Scientific) with excitation at 380 nm and emission at 460 nm. Caspase-3 activity was normalized to protein content and expressed as a percentage of that for control cells.

Determination of intracellular ROS. The level of intracellular ROS was expressed as a percentage of that for control cells. The determination of intracellular ROS was carried out as described previously.46 Values were normalized to the protein content and expressed as percentage of those for control cells. In addition, ROS levels were also detected by using flow cytometry analysis.

Assay for nuclear condensation and ultra-structural observation. Nuclear condensation was assessed by staining the nuclei with DAPI as described previously.47 In brief, the samples were then collected by centrifugation. The pooled cell pellets were fixed in 4% formaldehyde, washed with ice-cold PBS. Cells were then stained with 2.5 μg/ml DAPI solution for 10 min at room temperature, washed, resuspended and a fraction of the suspension was smeared on a slide. The slide was air dried, mounted with VECTASHIELD mounting medium, and examined under a DMI 4000 fluorescence microscope (Leica, Germany).

In addition, the transmission electron microscopy was utilized for analyzing the ultra-structural images of nucleus. In brief, treated cells were collected by centrifugation, samples were then fixed in 2.5% glutaraldehyde (Sigma-Aldrich, St Louis, MO, USA) for 2 h, washed in 0.1 M phosphate buffer (pH 7.4), post-fixed in 1% osmium tetroxide (Polysciences, Warrington, PA, USA) in 0.1 M phosphate buffer (pH 7.4) and subsequently dehydrated in increasing concentrations of alcohol. The samples were impregnated with proplene oxide (Merck-Schuchardt, Germany) and embedded in epoxy resin embedding media. After a light microscopic examination of semi-thin sections and staining with 2% (w/v) uranyl acetate and 1% (w/v) lead citrate, and then observed with a JEM-1200EXII (Jeol, Japan) transmission electron microscope.

Statistical analysis. All data are presented as the mean ± S.E.M. of at least three separate experiments. Statistical analysis was performed using one-way ANOVA followed by Dunnett’s multiple comparison test. Comparisons between two groups were analyzed using the Student’s t test. A P-value of less than 0.05 was considered to be statistically significant.

Conflict of interest

The authors declare no conflict of interest.

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