S-adenosyl-methionine decreases ethanol-induced apoptosis in primary hepatocyte cultures by a c-Jun N-terminal kinase activity-independent mechanism

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INTRODUCTION
Ethanol abuse increases the risk of developing liver damage, such as fatty liver, hepatitis, cirrhosis and the development of viral hepatitis and hepatocarcinomas[1-3]. Alcoholic liver disease (ALD) is a common health problem in Western countries, therefore the diagnostic evaluation and clinical management of ALD are major issues of concern. Clinical management depends on the extent of ALD and, although alcohol abstinence and supportive care are the classical treatments for ALD, the development of new therapies to improve ALD outcome are being examined. Among these new therapeutic agents are small c-jun N–terminal kinase (JNK) inhibitors, such as SP600125 and S-Adenosyl methionine (AdoMet), the principal biological methyl donor and also a glutathione (GSH) precursor in the liver[4-6]. However, an incomplete knowledge of the molecular mechanisms regulated by these agents in ALD development has complicated the validation of these new therapeutic approaches.
The JNK signaling pathway, a member of the mitogen-activated protein kinases family, regulates cell differentiation, proliferation, inflammation, cell survival and cell death in response to stress signals. Acute ethanol exposure induces JNK activation, promoting hepatocyte apoptosis, which has been strongly correlated with liver disease and postulated as the “nexus of liver injury and fibrosis.” Small molecule JNK inhibitors, such as SP600125, have been proved to exert a protective effect in reperfusion liver injury and are suggested as optional treatment for liver disease. Thus, the JNK signaling transduction pathway has emerged as an interesting element in liver disease development, opening the possibility that its modulation may represent an important approach for ALD treatment. Besides specific small molecule inhibitors, JNK activity may be modulated by other therapeutic agents, such as antioxidants. Indeed, in diverse aspects of cellular function, JNK is implicated as a redox stress cell sensor.

A possible hepatoprotective mechanism has been attributed to AdoMet effects on the regulation of cytosol and mitochondrial GSH levels. The up-regulation of GSH levels may have a down-regulatory effect on JNK signaling, since GSH acts as a key cell signaling modulator.

Although JNK has been identified as a pro-apoptotic signaling pathway involved in ALD development, the mechanism by which JNK induces programmed cell death remains far less clear, limiting the identification and validation of JNK-activity modulator agents as a potential ALD treatment. In this study, we determined the effect of the small JNK inhibitor SP600125 and AdoMet on the apoptotic pathway activated by ethanol and a possible modulating effect of AdoMet on JNK activity as part of its hepatoprotective mechanism. We found that JNK activation played a critical role in ethanol-induced apoptosis by means of its effect on Bid fragmentation, a key pro-apoptotic target of JNK. Even though AdoMet increased GSH concentration and protected against ethanol-induced apoptosis, we did not find an effect of AdoMet on JNK activity. These results strongly suggest that JNK activation induced by ethanol is independent of ROS generation, and that the AdoMet protective effects lie mainly at mitochondrial level.

MATERIALS AND METHODS

Materials

Anti-JNK-P and DeadEnd Colorimetric TUNEL System were obtained from Promega (Madison, WI). Antibodies against JNK and c-jun-P were purchased from Cell Signaling. c-jun, Bid, caspase 3 and cytochrome c antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Anti-actin monoclonal antibody was kindly provided by Dr. Manuel Hernández (CINVESTAV, IPN Mexico). JNK inhibitor SP600125 and caspase 8 inhibitor Z-IETD-fluoromethyl ketone were purchased from Calbiochem (San Diego, CA). Collagenase, ethanol, dimethyl sulfoxide (DMSO), AdoMet in p-toluene sulfonate salt form, 2', 7'-dichloro-dihydrofluorescein diacetate (DCFH-DA) and all other reagents were purchased from Sigma Chemical Company (St. Louis, MO). Electrophoresis reagents and Protein assay kit were from Bio-Rad (Richmond, Calif).

Animals

Male Fisher 344 rats with an average weight of 200 g were obtained from the Lab Animal Facility UPEAL, CINVESTAV. Animal care was performed according to the guidelines established by the Institutional Animal Care Committee.

Hepatocyte isolation and treatments

Rat hepatocytes were isolated from male Fisher rats (weighing 180-200 g) by collagenase perfusion as previously described. Viable hepatocytes were separated by Percoll gradient centrifugation at 1500 r/min for 2 min at 4 °C. Subsequently, cell viability was examined by trypan blue exclusion, and was over 90%. Hepatocytes were plated in Dulbecco’s modified Eagle’s medium (Gibco-BRL) supplemented with 100 mL/L bovine serum (Gibco-BRL). After 2 h incubation, the medium was changed to William’s E medium containing 100 mL/L bovine serum, 1.5 U/mL insulin, 20 nmol/L dexamethasone, 100 μg/mL streptomycin and 100 U/mL penicillin. Cells were cultured overnight at 37 °C in a humidified atmosphere containing 50 mL/L CO2. The day after plating, cells were washed with Hank’s balanced salt solution (HBSS), and incubated in William’s E medium (supplemented with 1 mL/L of fetal calf serum, 100 μg/mL streptomycin and 100 U/mL penicillin) with either 100 μmol/L SP600125, 25 μmol/L Z-IETD-FMK or 1 mL/L DMSO as a vehicle control for 2 h. Another set of cultures was pre-incubated with 4 mmol/L AdoMet for 1 h. Then 100 mmol/L ethanol was added and the dishes were sealed with parafilm to prevent evaporation. The cells were incubated for different periods of time at 37 °C before collection.

Cell viability assessment

Cell viability was determined using the MTT (methyl thiazole tetrazolium) assay as previously described. Briefly, at indicated times after pretreatments and ethanol exposure, media were removed and MTT solution (0.4 mg/mL in media, filter sterilized) was added and the cells were incubated for 2 h at 37°C. Plates were washed twice with PBS. The resultant formazan product was dissolved by addition of 500 μL DMSO to the plate. Optical density was measured at A595 nm and cell survival was expressed as percent of absorbance.

DNA gel electrophoresis

After different pretreatments and 24 h of ethanol exposure, hepatocytes (detached and attached) were harvested and washed in ice-cold PBS by centrifugation at 4°C at 1500 r/min for 5 min in an Eppendorf microcentrifuge. Approximately 8 × 105 cells were resuspended in lysis buffer (200 mmol/L EDTA; 100 mmol/L Tris, pH 8.0;...
8 g/L sodium lauryl sarcosine; 20 μg/mL DNase-free RNase) and incubated at 37 °C for 2 h, followed by addition of proteinase K (200 μg/mL) and another overnight incubation at 50 °C. DNA was electrophoresed on 18 g/L agarose gel at 35 V, stained with ethidium bromide and visualized under UV light[35].

**TUNEL assay**
The TUNEL assay was performed using a DeadEnd colorimetric apoptosis detection system kit. Rat hepatocytes were cultured on Falcon Chamber slides pre-coated with poly-L-lysine. After different pretreatments and 24 h ethanol exposure, slides were fixed by immersion in 40 g/L paraformaldehyde in PBS for 25 min at room temperature. Then, the TUNEL assay was performed as previously described[36] following the manufacturer's instructions. One hundred cells were counted in three randomly selected microscopic fields and cell apoptotic rate was expressed as a percentage of the total cells counted.

**Isolation of mitochondria**
Mitochondria were isolated from 4.8 x 10⁶ cells. Cells were washed twice with HBSS, scrapped and centrifuged for 10 min at 1500 r/min at 4 °C in an Eppendorf microcentrifuge. Cell pellets were resuspended in permeabilizing buffer (210 mmol/L D-mannitol, 10 mmol/L HEPES, 0.2 mmol/L EGTA, 50 mmol/L sucinate, and addition just before use of 70 mmol/L sarcarose, 1.5 g/L bovine serum albumin and 80 μg/mL digitonine, pH 7.2) and incubated at 4 °C for 20 min. Permeabilized cells were centrifuged at 170 r/min for 10 min at 4 °C. The supernatant was centrifuged at 13,000 g for 10 min at 4 °C, the second supernatant obtained contained the cytosolic fraction. The pellet resultant from the first centrifugation after the permeabilization step was incubated with 1 mL/L Triton X-100 in PBS for 20 min at 4 °C and centrifuged at 13,000 g for 10 min at 4 °C, thus the supernatant contained the mitochondrial fraction. Supernatants were precipitated with 50 g/L sulfosalicylic acid and resuspended in PBS[37]. Protein concentration in the supernatant was determined by the Lowry method.

**Immunoblot analysis**
Cellular protein was extracted at 4 °C in kinase lysis buffer (20 mmol/L Heps pH 8.0, 136 mmol/L NaCl, 1 mmol/L EDTA, 1 mmol/L EGTA, 10 mmol/L KCl, 2 mmol/L MgCl₂, 50 mmol/L sodium fluoride, 1 mmol/L phenylmethylsulfonyl fluoride, 1 mmol/L sodium orthovanadate, 2 g/L SDS) or Chaps buffer for samples to determine Bid and pro-caspase cleavage (50 mmol/L Tris-HCl, 2 mmol/L EDTA, 1 g/L Chaps, 1 mmol/L phenylmethylsulfonyl fluoride and Complete (BioRad) 25 x). Cell lysates were centrifuged at 12,000 g for 10 min at 4 °C. The supernatant protein concentration was determined by Lowry method. Whole cell extracts and cytosolic or mitochondrial fractions were resolved on SDS-PAGE and transferred onto nitrocellulose membrane. Blots were incubated overnight at 4°C with antibodies against JNK, phospho JNK, c-jun, phospho c-jun, Bid, caspase 3 (Bid antibody detected full length Bid and caspase 3 antibody recognized the precursor of pro-caspase 3 form) and cytochrome c. Then, blots were incubated with horseradish peroxidase-conjugated antibody and developed by chemiluminescence.

**Intracellular GSH content determination**
Hepatocytes were incubated under control conditions in William’s E medium only, or were pretreated with 4 mmol/L AdoMet for 1 h and then exposed to ethanol, or only exposed to ethanol. After the indicated times, cells were harvested and washed with HBSS buffer (pH 7.4). Cells (3x10⁶) were sonicated for 1 min in ice-cold 50 g/L sulfosalicylic acid, incubated for 20 min on ice and centrifuged at 12,000 g at 4 °C for 30 min. The resultant thiol extract was assayed by the method of Ellman with previously reported modifications[38,39]. The pellet obtained from the centrifugation was washed twice with 50 g/L sulfosalicylic acid and resuspended in 0.5 mol/L NaOH. Protein concentration was determined by the method of Lowry. The GSH content (including other eventual thiols) was expressed as nmol per mg of protein.

**Measurement of intracellular reactive oxygen species (ROS)**
The level of intracellular ROS was measured by the change in fluorescence resulting from oxidation of DCFDA. After treatment for indicated times, hepatocytes were washed with HBSS and incubated with 5 μmol/L DCFH-DA for 30 min at 37 °C. Hepatocytes were harvested and resuspended in HBSS, and then 5 g/L propidium iodide was added to detect dead cells. Intracellular ROS levels were measured on a Benckton Dickinson FACS Calibur. Ten thousand events were recorded for the analysis[40].

**Statistical analysis**
Data were analyzed using one-way ANOVA to determine differences between all independent groups. Differences between 2 groups were tested using two-tailed unpaired Student’s t-test. P<0.05 was considered statistically significant.

**RESULTS**

**Inhibition of ethanol-induced JNK activation by SP600125**
We first confirmed that 100 mmol/L ethanol exposure promoted JNK activation in our system. Ethanol induced JNK phosphorylation. We detected that the JNK1 basal activity was lower than JNK2, while JNK1 activity increment was higher than that for JNK2. Maximal JNK activation induced by ethanol was detected after 30 min and remained above control levels until 8 h after, as detected by c-jun phosphorylation, one of the most commonly measured parameters of JNK activity (Figure 1A-B). Next, we determined the effect of 100 μmol/L SP600125 on JNK activity. JNK activity significantly decreased in SP600125-pretreated hepatocytes compared to hepatocytes exposed to ethanol only (P<0.05), as determined by a decrease in c-jun phosphorylation (Figure 2A-B). The Western blot and densitometric analyses shown are representative of three individual experiments.
Inhibition of ethanol-induced apoptosis by SP600125 and AdoMet

Once we identified an effect of both ethanol and SP600125 on JNK activity, we determined the effect of SP600125 and AdoMet on the apoptosis process induced by ethanol. Ethanol-induced apoptosis was significantly reduced \( (P<0.05) \) in hepatocyte cultures pretreated with 100 \( \mu \)mol/L SP600125 or 4 mmol/L AdoMet, but not in hepatocytes pretreated with 1 g/L DMSO, the vehicle for SP600125, as shown by the DNA laddering assay (Figure 3A) or TUNEL assay (Figure 3B). A 24-h exposure to 100 mmol/L ethanol produced approximately 47% cell death in non-pretreated hepatocytes as compared to 23% and 25% cell death observed in SP600125- and AdoMet-pretreated cultures, respectively. Cells pretreated with DMSO but not stimulated with ethanol showed 5% cell death (Figure 1C).

Protection of SP600125 and AdoMet on ethanol hepatocyte injury

In order to determine whether or not the survival of hepatocytes exposed to 100 mmol/L ethanol was modulated by SP600125 and AdoMet, we determined survival rates after different periods of 100 mmol/L ethanol exposure. Cell metabolic activity, determined by the MTT assay, significantly decreased after 12 h in cells only exposed to ethanol \( (P<0.05) \) compared to the cells pre-incubated with AdoMet, SP600125 or incubated under control conditions. Cell metabolic activity, measured as the percentage of tetrazolium salt reduction, decreased from 75% observed in pretreated hepatocytes to 40% observed in hepatocytes exposed to ethanol only (Figure 4).

AdoMet effect on GSH intracellular levels from ethanol-incubated hepatocytes

The JNK signaling pathway activation may be modulated by antioxidants. AdoMet, a GSH precursor in liver, prevented ethanol-induced apoptosis as aforementioned. We tested if, through an antioxidant effect, AdoMet may be related to an inhibitory action of JNK activity as a possible hepatoprotective mechanism. We first determined the effect of 4 mmol/L AdoMet pretreatment on GSH intracellular levels in ethanol-incubated hepatocytes. Pre-incubation with 4 mmol/L AdoMet significantly prevented GSH decrease induced by the exposure of cultured hepatocytes to 100 mmol/L ethanol \( (P<0.05) \), and moreover, maintained GSH levels above those found in control hepatocyte cultures. A significant increase in GSH concentra-
tion was observed between 8 and 12 h in hepatocytes pretreated with AdoMet compared to hepatocytes incubated in control medium (Figure 5) \( (P < 0.05) \). Furthermore, ROS generation in hepatocytes pretreated with 4 mmol/L AdoMet and then exposed to ethanol was maintained as in control cells in comparison to hepatocyte cultures exposed to ethanol only, in which a significant ROS generation was observed \( (P < 0.05) \).

**Figure 3** Effect of SP600125 and AdoMet in apoptosis induced by ethanol. A: DNA ladder assay; B: 40x TUNEL assay; C: data from TUNEL assay; a: control; b: ethanol; c: AdoMet+ ethanol; d: SP600125 + ethanol. The data represent mean ±SEM of \( n = 3 \). *indicates a significant difference \( (P < 0.05) \) between SP600125 or AdoMet-pretreated cells and cells only pretreated with DMSO and exposed to ethanol.

**Figure 4** Protective effect of SP600125 and AdoMet on ethanol hepatocyte injury. Data represent percentage of MTT reduction in mean±SEM of \( n = 3 \). *represents a significant difference \( (P < 0.05) \) in MTT reduction between pretreated and non-pretreated cells.

**Figure 5** Prevention of GSH decrease in hepatocyte cultures exposed to ethanol by AdoMet. Data represent percentage change of ROS levels in mean±SEM of \( n = 3 \). *represents a significant difference \( (P < 0.05) \) in ROS generation between pretreated cells and cells exposed to ethanol only.

**Effect of AdoMet on JNK activity**

Once we determined that AdoMet had antioxidant effects, we tested if AdoMet exerted an inhibitory action on JNK activity. We first measured AdoMet effect on c-jun phosphorylation. AdoMet pretreatment slightly decreased JNK activity induced by ethanol treatment by 25.4% with respect to the cultures exposed to ethanol only, contrasting with the effect of the selective JNK inhibitor which produced a c-jun phosphorylation inhibition of approximately 68.8% with respect to ethanol- treated samples. So, the decrease on c-jun phosphorylation induced by AdoMet was
Effects of SP600125 and AdoMet on Bid fragmentation, another JNK target

We next determined the effects of SP600125 and AdoMet on Bid cleavage. Bid is a proapoptotic member of the Bcl-2 family, which may represent another important JNK target. We observed that 100 mmol/L ethanol hepatocyte exposure promoted Bid cleavage between 4 h and 24 h. SP600125 hepatocyte pretreatment significantly prevented Bid fragmentation compared to cells stimulated with ethanol only between 4 h to 24 h (P < 0.05). Bid has been reported as a caspase 8 target. Indeed, we found that preincubation with 25 μmol/L Z-IETD-fmk, a selective caspase 8 inhibitor, significantly decreased Bid fragmentation induced by ethanol (Figure 8) (P < 0.05). These results suggested that Bid cleavage induced by ethanol not only depends on JNK, but also on caspase 8 activity. Nevertheless, even if AdoMet exerted antioxidant properties, it did not prevent Bid fragmentation induced by ethanol. Western blot and densitometric analyses shown are representative of three independent experiments.

Decrease in ethanol-induced cytochrome c release and caspase 3 activation by SP600125 and AdoMet

Apoptosis induced by 100 mmol/L ethanol involves Bid fragmentation, which is responsible for cytochrome c release with the subsequent activation of caspase 3. We determined the effect of SP600125 and AdoMet on these apoptotic components. Pretreatment of hepatocyte with SP600125 prevented Bid cleavage induced by ethanol and decreased cytochrome c release from mitochondria between 8 h and 24 h after ethanol exposure. Even though AdoMet did not have an effect on Bid fragmentation, it decreased cytochrome c release from mitochondria (Figure 9). Cytochrome c release contributes to pro-caspase 3 fragmentation, so we observed that cytochrome c release prevention by SP600125 and AdoMet also decreased pro-caspase 3 fragmentation (Figure 10). These results indicated that the hepatoprotective effects of SP600125 and AdoMet converge at mitochondrial level, but AdoMet antiapoptotic action mainly lies in an inhibition of the mitochondrial apoptotic pathway.
This study points out that modulation of the JNK signaling transduction pathway by selective JNK inhibitors and mitochondrial function preservation by AdoMet are among several key factors that must be considered to develop successful therapies for ALD treatment. We determined the role of JNK activity in the apoptotic process induced by ethanol in primary hepatocyte cultures and the possible modulator effect of JNK activity by AdoMet, an agent increasingly used in ALD treatment.

JNK activity is related to proapoptotic responses, but it is also linked to antiapoptotic signals and cell proliferation\(^{14,15}\), thus the functions and components of this signaling cascade still remain controversial. We exposed hepatocytes to 100 mmol/L ethanol, considering previous reports which indicate that ethanol levels in the peripheral blood of some alcoholic patients have been exceeded 100 mmol/L\(^ {14}\). In the present study, the exposure of primary hepatocyte cultures to 100 mmol/L ethanol promoted JNK activation. We detected that JNK1 basal activity was lower than JNK2 basal activity, while the JNK1 activity increment was higher than JNK2. These findings are in agreement with previous reports\(^ {9,14}\) and strongly suggest that JNK1 activity is related to the proapoptotic function of this signaling pathway activation by ethanol. We determined ethanol JNK activation by measuring c-jun phosphorylation, the most common parameter of JNK phosphorylation, the most common parameter of JNK activation. We detected that JNK1 basal activity was significantly decreased by 100 μmol/L SP600125, a concentration previously reported for the selective inhibition of JNK\(^ {15}\). Indeed, c-jun phosphorylation was significantly decreased by 100 μmol/L SP600125 in comparison with cells pretreated with 1 g/L DMSO only, as a vehicle control. We measured a decrease in c-jun and not in JNK phosphorylation because previous reports have indicated that SP600125 specifically inhibits c-jun phosphorylation and does not down-regulate p-JNK levels\(^ {14,15}\). In our study, selective JNK inhibition by SP600125 allowed us to confirm that JNK functioned as a proapoptotic signaling pathway in hepatocyte cultures exposed to 100 mmol/L ethanol for 24 h, since in SP600125-pretreated hepatocyte cultures, DNA ladder and positive TUNEL cells significantly decreased. Our results, in agreement with previous reports\(^ {9,14}\), indicated that JNK might be a key pathway involved in apoptosis induction by ethanol. However, other signaling events must be involved in the apoptotic pathway induced by ethanol, since apoptotic cells in SP600125-pretreated hepatocytes were not completely abolished.
suggesting that JNK activity contributes to but is not the only factor required for the apoptotic process induced by ethanol. MAPK p38 is also a key candidate in this process, since there are several reports indicating that TNF-alpha, which plays a critical role in ALD development, induces p38 activation\[40]. Recently, Bid, a pro-apoptotic member of the bcl-2 family, has been reported as a JNK target in different systems\[46,47]. We observed that the incubation of primary hepatocyte cultures with 100 mmol/L ethanol induced Bid cleavage, which was decreased by SP600125. These results strongly suggested Bid as a key target of the pro-apoptotic JNK activity induced by ethanol. Even though maximal JNK activity, measured by c-jun phosphorylation, was observed between 30 min and 4 h after ethanol treatment, Bid fragmentation was not observed until 8 h after ethanol exposure, when c-jun phosphorylation was slightly above control. This slow kinetic of JNK-stimulated Bid cleavage may indicate that several steps separate JNK activation from Bid fragmentation\[48]. Since JNK is not a protease, it is likely that one of these steps corresponds to a protease induction by JNK\[49,50]. Thus, the exact target of JNK involved in Bid fragmentation is yet to be established\[51]. Bid cleavage was also decreased when cultures were exposed to Z-IEETD-fmk, a selective caspase 8 inhibitor, indeed it has been reported that Bid is a caspase 8 target\[52]. Fas-mediated apoptosis has been related to FasL induction by c-jun activation, and downstream to caspase 8 and Bid fragmentation. Our result suggests that Bid fragmentation in the apoptosis process induced by ethanol lies at least in two distinct pathways, and not only depends on JNK activity. In accordance with this result, a previous report indicates that Bid fragmentation by JNK does not depend on transcription, and in addition, SP600125 does not affect Fas-mediated apoptosis\[53]. JNK and other kinases that regulate JNK activity are known to be responsive to redox changes\[27,55\], and oxidative stress is critically involved in ALD development\[28,29,30,31,33,34\]. We observed that treatment of hepatocyte culture with 100 mmol/L ethanol induced a peak of ROS generation between 6 h and 8 h after ethanol exposure, which coincided with a significant decrease in GSH levels. These redox alterations may be resulted from two of the main components of alcohol metabolism, alcohol dehydrogenase (ADH) and cytochrome P450 2E1, since it has been reported under similar culture conditions that 4-methyl pyrazole, an ADH inhibitor, led to the reduction of H₂O₂ generation\[21,53\]. Besides, different non-enzymatic free radical pathways and the mitochondrial dysfunction after the reduction of GSH content are involved in ROS generation induced by ethanol. Therefore, JNK activation in response to ethanol may be related to the oxidative metabolism of ethanol in the liver and to the resultant alterations in intracellular redox state, primarily reflected by GSH changes. AdoMet, an increasingly used agent for liver disease treatment, has wide physiological and pharmacological actions, including its effect on GSH metabolism. However, its hepatoprotective mechanisms have not been completely elucidated\[27,28\]. We observed that AdoMet, as SP600125, significantly reduced ethanol-induced apoptosis. AdoMet has been identified as a GSH precursor, a major cell antioxidant, and JNK activity has been reported to be down-regulated by antioxid-
dant agents. Thus, our study suggested that AdoMet may down-regulate JNK activity as part of its hepatoprotective mechanism through an antioxidant effect. We found that 4 mmol/L AdoMet was able to prevent GSH decrease induced by ethanol in primary hepatocyte cultures and even maintained intracellular GSH levels above those found in non-treated cells. Furthermore, AdoMet pretreatment also diminished ethanol-induced ROS generation. These results indicated that AdoMet might be related to the repletion of GSH levels. There are contradictory results about the effect of AdoMet on GSH levels; while a previous study found no effect of AdoMet on GSH levels\[27\], present results are in agreement with others which reported an increment of GSH levels resulted from AdoMet pretreatment\[29,30\]. Even though AdoMet exerted a redox state regulating effect, surprisingly, it only decreased c-jun phosphorylation by 25.5% in comparison to the inhibition induced by SP600125, which corresponded to 62.8%. We also tested the effect of AdoMet on Bid fragmentation and did not observe an inhibitory effect of AdoMet until 24 h after ethanol exposure. At this time, JNK activity was below control levels, so it is possible that this decrease was not related to JNK activity. These results indicated that even though AdoMet exerted an antioxidant action, it did not have a significant effect on JNK activity, at least on c-jun or Bid, the measured JNK targets. Thus, we suggest that JNK activation may be independent of oxidative stress, in accordance with previously reported results in which cell pretreatment with N-acetyl-cysteine failed to prevent JNK activation\[40\]. Other reports suggest that JNK activation may be independent of ethanol metabolism, since ethanol activates JNK in HepG2 cells which do not express detectable levels of Cyp2E1 and express low levels of alcohol dehydrogenase activity\[56\]. However, further studies are required to determine the regulatory mechanisms of JNK activity independent of ROS generation. Even though AdoMet did not have a significant effect on JNK activity as it was measured by its action on c-jun and Bid, it preserved the mitochondrial metabolic activity as it was measured by the MTT assay. In hepatocytes exposed to ethanol only, the mitochondrial function was significantly reduced compared to AdoMet-pretreated hepatocytes. Downstream AdoMet prevented cytochrome c release and pro-caspase 3 fragmentation. In agreement with previous reports which indicated that AdoMet maintained mitochondrial membrane stability and corrected the reduction of GSH content both in cytosol and mitochondria\[29,33,34\], these results suggest that the AdoMet protective effect mainly lies at mitochondrial level.

In summary, our results point out that ALD is a complex and multifactorial process, and that the mechanisms involved in ALD development must be thoroughly explored for the design of successful therapeutic strategies. We found that AdoMet, although protected against ethanol apoptosis induction and exerted an antioxidant effect, did not show any inhibitory action on Bid fragmentation, a key proapoptotic target of JNK. This finding is in agreement with the suggestion that the mechanisms which lead to JNK activation are independent of ROS generation. We conclude that the JNK signaling pathway modulation by selective inhibitors.
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