In vitro Chemo-protective effect of bioactive peptide lunasin against oxidative stress in human HepG2 cells

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Running title: Lunasin protects HepG2 from oxidative stress

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ABSTRACT

Lunasin is a peptide with proven properties against cancer and cardiovascular diseases. Relevant amounts of lunasin have been found in liver of rats fed lunasin-enriched diets, indicating its potential bioactive effect in this tissue. This study investigated the stability of lunasin in human liver HepG2 cells, and its chemoprotective effect against oxidative stress induced by tert-butyldihydroperoxide. Cell viability and biomarkers of redox status were evaluated. Pre-treatment of cells with lunasin (0.5-10 μM) significantly prevented the increased reactive oxygen species (ROS) generation (122% compared to 190% in stressed cells), and glutathione peroxidase and catalase activities, as well as the depletion of reduced glutathione. By restraining ROS overproduction, lunasin evoked a decline in carbonyl groups, and a significant recovery from cell death by apoptosis. These findings suggest that lunasin, at physiological concentrations, might confer a significant chemoprotection against oxidative stress-associated liver disorders. In addition, fragments released after hydrolysis of lunasin by cell enzymes might contribute on the observed antioxidant effects.

Keywords: antioxidant defences, biomarkers for oxidative stress, dietary antioxidants, peptide lunasin
1. INTRODUCTION

Cells are naturally provided with an extensive array of protective enzymatic and non-enzymatic antioxidants that counteract the potentially injurious oxidizing agents. But even this multifunctional protective system cannot completely prevent the deleterious effects of reactive oxygen species (ROS), and consequently, molecules damaged by oxidation accumulate in cells. Large amounts of ROS have been shown to participate in the pathogenesis of several human degenerative diseases, including inflammation, cardiovascular and neurodegenerative disorders, and cancer (Ramos, 2008). Restoration or activation of improperly working or repressed antioxidant machinery as well as suppression of abnormally amplified inflammatory signalling can provide important strategies for chemoprevention. Therefore, determination of anti-inflammatory and/or antioxidant properties has been proposed as a good indicator for screening anti-cancer agents (Federico, Morgillo, Tuccillo, Clardiello, & Loguercio, 2007).

There is substantial evidence that antioxidant food components have a protective role against oxidative stress-induced atherosclerosis, degenerative and age-related diseases, cancer and aging.¹ Food-derived peptides are promising natural antioxidants without marked adverse effects. In addition to their potential as safer alternatives to synthetic antioxidants used to avoid or retard oxidation reactions in foods, antioxidant peptides can also act reducing the risk of numerous oxidative stress-associated disorders (Meisel, 2004). Lunasin is a 43-amino acid peptide identified in soybean and other seeds and plants which chemopreventive properties have been recently reviewed (Hernández-Ledesma, Hsieh, & de Lumen, 2013). This peptide has demonstrated, by using cell cultures and animal models, to act as anticarcinogenic agent against skin, prostate, colon, and breast cancer (Hernández-Ledesma, de Lumen, & Hsieh, 2013). Recently, lunasin has been commercialized in the US by its benefits on cardiovascular system through reduction of low density lipoprotein cholesterol concentration (Gálvez, 2012).
Moreover, its promising anti-inflammatory and antioxidant activities reported in the recent years might contribute on lunasin chemoprotective action. Lunasin has been shown to inhibit inflammation in cultured RAW 264.7 macrophages through suppression of nuclear factor (NF)-κB pathway (González de Mejia, & Dia, 2009; Hernández-Ledesma, Hsieh, & de Lumen, 2009a; Cam, Sivaguru, & González de Mejia, 2013). Additionally, in vitro assays have revealed the ability of this peptide to scavenge peroxyl radicals, and to block Fenton reaction by chelating iron ferrous ions, protecting DNA from oxidative damage (Hernández-Ledesma et al., 2009a; Jeong, de Lumen, & Jeong, 2010; García-Nebot, Recio, & Hernández-Ledesma, 2014).

Studies on bioavailability carried out in mice and rats have demonstrated that, after its oral ingestion, lunasin appears in an intact and active form in different organs and tissues, such as blood, liver, and kidney, among others (Jeong, Lee, Jeong, Park, Cheong, & de Lumen, 2009; Hsieh, Hernández-Ledesma, Jeong, Park, & de Lumen, 2010). In humans, presence of lunasin has been also reported in plasma, indicating that this peptide might reach different target tissues, and exert its biological activity (Dia, Torres, de Lumen, Erdman, & González de Mejia, 2009).

The liver is particularly susceptible to toxic and oxidative insults since the portal vein brings blood to this organ after intestinal absorption. Therefore, studies dealing with the effects of chemopreventive compounds at a cellular level in cultured hepatic cells are essential. Human HepG2, a well differentiated transformed cell line, is a reliable model for cultured hepatocyte-type cells used for biochemical, pharmacological and nutritional studies since it retain hepatocyte morphology and most of its functionality in culture (Alía, Ramos, Mateos, Bravo, & Goya, 2006; Mateos, Goya, & Bravo, 2006).

The aims of the study were to evaluate the stability of lunasin in human liver HepG2 cells as a model for cultured hepatocytes, and to investigate the potential chemo-protective effect
of this peptide against oxidative stress chemically induced by a potent pro-oxidant, tert-butyl hydroperoxide (t-BOOH). Cell integrity and several biomarkers of oxidative damage were evaluated to estimate the effect of lunasin on cell survival and on the response of the antioxidant defence systems of HepG2 cells to t-BOOH.

2. MATERIALS AND METHODS

2.1. Reagents

Peptide lunasin (>95% of purity) was synthesized by Chengdu KaiJie Biopharm Co., Ltd (Chengdu, Sichuan, P. R. China). t-BOOH, glutathione reductase, reduced glutathione (GSH), nicotine adenine dinucleotide phosphate reduced salt (NADPH), o-phthalaldehyde (OPT), dichlorofluorescin (DCFH), dinitrophenylhydrazine (DNPH), trifluoroacetic acid (TFA), ethylenediaminetetraacetic acid (EDTA), β-mercaptoethanol, gentamicin, penicillin G, streptomycin, Triton-X100, 4-(2-hydroxyethyl)piperazine-1-ethanesulfonic acid (HEPES) and dithiothreitol (DTT) were purchased from Sigma Chemical Co. (Madrid, Spain). Sodium dodecyl sulphate (SDS) was from Panreac Química (Madrid, Spain). N-acetyl-Asp-Glu-Val-Asp-7-amino-4-methylcoumarin (Ac-DEVD-AMC) was from BD Pharmigen (Madrid, Spain). Bradford reagent was from BioRad Laboratories S.A. (Madrid, Spain). The rest of chemicals used were of HPLC grade.

2.2. Cell culture

Human HepG2 cells were grown in a humidified incubator containing 5% CO₂ and 95% air at 37°C. They were grown in Dulbecco's Modified Eagle Medium (DMEM)-F12 medium from Biowhitaker (Lonza, Madrid, Spain), supplemented with 2.5% (v/v) Biowhitaker foetal bovine serum (FBS) and 50 mg/L of each of the following antibiotics: gentamicin, penicillin
G, and streptomycin. Cells were changed to FBS-free medium the day before the assay (Mateos et al., 2006).

2.3. Cell treatment conditions

Cells were incubated for 20 h with peptide lunasin dissolved in FBS-free DMEM-F12 at final concentrations ranging from 0.5 to 10 μM. To evaluate both direct and protective effects against oxidative stress, the incubation period was followed by a 3 h treatment with culture medium (direct effect) or oxidant chemical t-BOOH (400 μM). Crystal violet (CV) staining, GSH concentration, ROS generation and glutathione peroxidase (GPx) and catalase (CAT) activities were evaluated in both direct and protective experiments. Besides, the protective effect of lunasin against t-BOOH-induced oxidative damage to proteins and apoptotic cellular signals was evaluated by carbonyl groups and caspase-3 assays, respectively.

2.3.1. Crystal violet assay

Cell viability was evaluated following the CV assay described by Granado-Serrano and co-workers (Granado-Serrano, Martín, Izquierdo-Pulido, Goya, Bravo, & Ramos, 2007). HepG2 cells were seeded in 96-well plates (1 × 10⁴ cells per well) and incubated overnight. Then, cells were treated with lunasin for 20 h as described above, washed with PBS, and incubated with CV (0.2% in ethanol) for 20 min at room temperature. Finally, cell lysis was carried out with 1% SDS, and the absorbance was read at 570 nm in a microplate reader (FL600, Bio-Tek, Winooski, VT, USA). Results were pooled from different plates to obtain average of n = 12, and presented as percentage of viable cells compared to control, considered as 100%.
2.3.2. Morphological analysis

HepG2 cells were exposed to increasing concentrations (0.5-10 μM) of lunasin for 20 h, then treated with DMEM (controls) or DMEM supplemented with t-BOOH for 3 h, and cell images were taken using an inverted phase contrast microscope at 20 × magnification.

2.3.3. Determination of intracellular ROS levels

Intracellular ROS levels were quantified following the method described by Alía and co-workers, using DCFH as fluorescent probe (Alía, Ramos, Mateos, Bravo, & Goya, 2005). HepG2 were cultured in 24-wells multiwell plates (2 × 10⁵ cells per well), incubated overnight, and then, treated with lunasin as described above. After 19.5 h incubation with the peptide, a solution of DCFH was added to each well (5 μM final concentration), and cells were incubated in the dark for 30 min at 37°C. Afterwards, cells were washed with PBS, incubated with culture medium (direct effects), or subjected to chemical-induced oxidative stress with t-BOOH (protective effects). The production of intracellular ROS was followed over 3 h measuring the fluorescence intensity at λ(excitation) and λ(emission) of 485 nm and 530 nm, respectively, in a microplate reader (FL600, Bio-Tek). The results were pooled from different plates to obtain average of n = 8, and expressed as percentage of the control (cells in a basal state), considered as 100%.

2.3.4. Quantification of concentration of GSH and determination of GPx and CAT activity

Cells were seeded onto 100-mm Petri dishes, incubated overnight, treated with lunasin (for both direct and protective assays), exposed to t-BOOH (for protective assays), and then, collected following the methodology described by Quéguineur and co-workers (Quéguineur, Goya, Ramos, Martín, Mateos, & Bravo, 2012). The obtained supernatants, corresponding to the cellular content of HepG2 cells, were subjected to the determination of the concentration
of GSH and the activity of GPx and CAT enzymes. The content of GSH was evaluated by a fluorometric assay as previously described (Quéguineur et al., 2012). Briefly, 50 µL of each sample were transferred in triplicate to a 96-multiwell plate, and the reaction mixture containing 15 µL of 1 M NaOH, 175 µL of PBS/EDTA, and 10 µL of a solution of OPT (10 mg/mL) was added. After 20 min incubation, the fluorescence was read (λ_{excitation}/λ_{emission} = 340/460 nm) in a microplate reader (FL600, Bio-Tek). Results were pooled from different plates to obtain average of n = 6, and interpreted considering those of a standard GSH curve similarly prepared within each of the experiments run.

GPx catalyses the oxidation of GSH to oxidized glutathione, using t-BOOH as a substrate, reaction coupled to the decomposition of NADPH to β-nicotinamide adenine dinucleotide by glutathione reductase. Thus, disappearance of NADPH reliably estimates GPx activity, which can be measured by following the decrease in absorbance at 340 nm (Alía et al., 2006). CAT activity was determined by following the breakdown of the peroxide H$_2$O$_2$ to H$_2$O, monitored as a decrease in absorbance at 240 nm (Granado-Serrano et al., 2007). Results for both GPx and CAT activities were pooled from different plates to obtain average of n = 6, and 4, respectively, and referred to the total protein concentration of the cytosolic samples, measured by the Bradford reagent (Bio-Rad).

2.3.5. Evaluation of carbonyl groups

Oxidative damage to proteins by reactive species, particularly ROS, was evaluated by measuring the content of carbonyl groups in cell supernatants according to the method of Richert and co-workers (Richert, Wehr, Stadtman, & Levine, 2002). Absorbance was measured at 360 nm, and carbonyl content results (experiments run to obtain an average of n = 4) were expressed as nmol of carbonyl groups per mg of total protein, using an extinction coefficient of 22000 nmol/L/cm.
2.3.6. Determination of caspase-3 activity

Caspase-3 activity was measured according to the fluorometric assay previously described (Herrera et al., 2001). After treatment with lunasin for 20 h and chemical induction with t-BOOH for 3 h, cells were collected, and lysed in a buffer containing 5 mM Tris, 20 mM EDTA, and 0.5% Triton-X100. Then, the reaction mix containing 30 μg cell protein, 20 mM HEPES, 10% glycerol, 2 mM DTT, and 20 μM Ac-DEVD-AMC was incubated in the dark for 2 h. Fluorescence was measured at \( \lambda_{\text{excitation}} \) and \( \lambda_{\text{emission}} \) of 380 nm and 440 nm, respectively, in a microplate reader (FL600, Bio-Tek), and enzymatic activity results (experiments run to obtain an average of \( n = 4 \)) were expressed as units of caspase-3 per μg of total protein.

2.4. Stability of lunasin

HepG2 cells were plated in 60 mm-diameter plates at a density of 1.5 × 10^6 cells per plate, and incubated at 37°C overnight. Then, cells were treated with 10 μM lunasin dissolved in FBS free DMEM-F12, and incubated at 37°C for 20 h, taking aliquots of the medium at 0, 2, 6, 12, and 20 h of incubation. These aliquots were subjected to liquid chromatography (HPLC) coupled to tandem mass spectrometry (HPLC-MS/MS) on an Agilent 1100 HPLC System (Agilent Technologies, Waldbronn, Germany) connected on-line to an Esquire 3000 ion trap (Bruker Daltonik GmbH, Bremen, Germany), and equipped with an electrospray ionization source as previously described (Contreras, Gómez-Sala, Martín-Álvarez, Amigo, Ramos, & Recio, 2010). The column used was a Mediterranea Sea 18 (150 x 2.1 mm, Teknokroma, Barcelona, Spain), the injection volume was 50 μL, and the flow was set at 0.2 mL/min. Peptides were eluted with a linear gradient of solvent B (acetonitrile:TFA 1000:0.27 v/v) in A (water:TFA 1000:0.37 v/v) going from 0% to 45% in 130 min. Data obtained were processed and transformed to spectra representing mass values using the Data Analysis
program (version 4.0, Bruker Daltonik). To process the MS/MS spectra and to perform peptide sequencing BioTools (version 3.1, Bruker Daltonik) was used.

2.5. Statistics

Data were analyzed by a one-way ANOVA followed by the Bonferroni Multiple Comparison test, and expressed as the mean ± standard variation (SD). GraphPad Prism 5.0 software (San Diego, CA, USA) was used to perform statistical analyses. Differences with a $P$ value < 0.05 (*, #), $P$ value < 0.01 (**, ##) or $P$ value < 0.001 (***, ###) were considered significant.

3. RESULTS AND DISCUSSION

3.1. Chemoprotective effects of lunasin on cell viability and redox status

In order to evaluate the effect of lunasin at physiological level, in the present study, a range of concentrations between 0.5 to 10 μM was selected. Previous studies on lunasin’s bioavailability have demonstrated that, because of the protection against gastric and pancreatic enzymes exerted by naturally occurring protease inhibitors such as the Bowman-Birk inhibitor (BBI), a high percentage of daily ingested peptide remains intact during its passage through gastrointestinal tract, reaching target organs and tissues in an active form (Hsieh et al., 2010). The presence of lunasin and BBI has been confirmed in different soybean products, such as soymilk, tofu, soybean cake and fermented soybean products (Hernández-Ledesma, Hsieh, & de Lumen, 2009b). Daily consumption of 25 g of soy protein, recommended by Food and Drug Administration (FDA) to reduce coronary disease (FDA, 1999), supplies lunasin in quantity ranged from 110 mg (21.9 μmol) to 1760 mg (350.2 μmol). Selected concentrations in our study were not far from reality since steady-state concentrations of 99.3 μg/g tissue were isolated from liver of rats fed lunasin-enriched rye
diets for 4 weeks, and lunasin extracted from those livers has been demonstrated to be active (Jeong et al., 2009).

Before testing the chemo-protective effect of lunasin, it was important to evaluate the effect of this peptide per se, in basal conditions, ensuring that no direct damage is caused to the cell by the compound. Thus, the direct effect of lunasin on HepG2 cells viability was evaluated. The CV assay, based on the growth rate reduction reflected by the colorimetric determination of the stained cells, was used to determine the cell viability. As shown in Figure 1A, treatment of HepG2 cells with lunasin for 20 h evoked no decreases in CV staining, indicating that the concentrations selected for the study (0.5-10 μM) did not damage cell integrity during the period of incubation. Higher concentrations of lunasin were also assessed, observing that the percentage of viable cells were not affected by 25 μM lunasin (111.20% ± 4.80) and 50 μM lunasin (111.07% ± 6.19).

Because of its connection to the gastrointestinal tract and its unique metabolic activity, the liver is one of the main body targets of the toxicity of drugs, xenobiotics, environmental contaminants, as well as oxidative stress (Jaeschke, Gores, Cederbaum, Hinson, Pessayre, & Lemasters, 2002). The cell damage caused by ROS and other reactive species plays a crucial role in the induction and progression of several liver diseases such as hepatocarcinoma, viral and alcoholic hepatitis, and non-alcoholic steatosis. Consequently, there is an increasing interest in new therapeutic agents protecting liver from such oxidative damage, with natural antioxidants being considered one of the most effective alternatives (Vitaglione, Morisco, Caporaso, & Fogliano, 2004). t-BOOH, a short-chain analog of lipid peroxide, is often used to induce acute oxidative stress in different *in vitro* and *in vivo* systems, and to evaluate the protective effects of antioxidants (Mersch-Sundermann, Knasmuller, Wu, Darroudi, & Kassie, 2004). In the case of hepatocytes, these cells metabolize t-BOOH to toxic peroxyl and alkoxy radicals, initiating peroxidation of macromolecules, and thus, affecting the cell integrity and
leading to chemical-induced hepatic oxidative damage. Therefore, treatment of HepG2 cells with t-BOOH is an excellent model of oxidative stress in cell culture systems (Alía et al., 2005). To evaluate whether lunasin protects HepG2 from oxidative stress induced by t-BOOH, we first determined its effects on the cell viability. As shown in Figure 1A, the percentage of viable cells in HepG2 induced by 400 μM t-BOOH was 67.5% (compared to non-stressed cells), indicating that this chemical showed remarkable cytotoxicity ($P < 0.001$) on these cells. However, incubation of the cells with 0.5-10 μM lunasin for 20 h, prior to chemical oxidant treatment, significantly restored cell viability up to 99.8% ($P < 0.001$), although dose-dependent effects were not observed.

As shown in Figure 2A and 2B, HepG2 cells treated with t-BOOH showed reduction in cell numbers and loss of cell-to-cell contact. These morphological changes in t-BOOH-induced HepG2 cells were attenuated by pre-treatment with lunasin at 0.5 μM (Figure 2C), and 5 μM (Figure 2D).

As shown in Figure 2A, 2B, morphological changes in HepG2 cells due to the toxic effect of t-BOOH were clearly visible. However, such changes were reduced by lunasin at 0.5 μM (Figure 2C), and 5 μM (Figure 2D).

In order to understand whether the cytoprotective effect of lunasin might be attributed to the reduction of oxidative stress, the intracellular ROS generation was evaluated in HepG2 exposed to t-BOOH with and without pre-treatment with lunasin for 20 h. As shown in Figure 1B, the level of intracellular ROS in HepG2 cells treated with t-BOOH alone was 190.0% compared to the non-treated cells (considered as 100%), indicating that 400 μM t-BOOH had a strong effect on ROS generation. When the cells were pre-treated with lunasin, intracellular ROS levels were significantly decreased up to 122% (compared to control) ($P < 0.001$), but no dose-dependent activity was observed. These results suggest that ROS generated during the period of oxidative stress were more efficiently quenched in cells pre-treated 20 h with
lunasin, which could be a first explanation for the reduced cell damage and death shown. The intracellular ROS levels of non-stressed cells were also decreased by treatment with lunasin, reaching 63.6% (compared to control cells) when 5 μM lunasin was used (Figure 1B). These findings confirm the ability of this peptide as an effective scavenger of ROS in cell cultures. Previous studies have demonstrated that lunasin is a potent antioxidant in different in vitro assays, including reducing power, and ABTS, peroxyl and superoxide radical scavenging activity (Hernández-Ledesma et al., 2009a; Jeong et al., 2010; García-Nebot et al., 2014).

Lunasin at concentrations higher than 1 μM significantly increased the cytosolic levels of GSH ($P < 0.05$) (Figure 1C). GSH is a tripeptide found in all mammalian cells and considered the main non-enzymatic antioxidant defense within the cell. It appears in high concentration in the liver where plays a crucial role protecting against oxidative stress through elimination of toxic ROS, and mitigation of macromolecules peroxidation and cell injury (Mari, Morales, Colell, García-Ruiz, & Fernández-Checa, 2009). It is usually assumed that enhanced levels of GSH prepare the hepatocytes against a potential oxidative insult, whereas its depletion reflects an intracellular oxidation state (Alía et al., 2005). The effect of lunasin on GSH levels, together the decreased ROS generation, reflects a diminished intracellular oxidation which could be expected to place the cell in favourable conditions to face a potential oxidative insult. In addition to its effect as inducer of ROS generation, t-BOOH also provokes GSH depletion (Goya, Mateos, & Bravo, 2007). GSH is a tripeptide found in all mammalian cells and considered the main non-enzymatic antioxidant defense within the cell. It appears in high concentration in the liver where plays a crucial role protecting against oxidative stress through elimination of toxic ROS, and mitigation of macromolecules peroxidation and cell injury (Mari, Morales, Colell, García-Ruiz, & Fernández-Checa, 2009). It is usually assumed that enhanced levels of GSH prepare the hepatocytes against a potential oxidative insult, whereas its depletion reflects an intracellular oxidation state (Alía et al., 2005). Lunasin at
concentrations higher than 1 μM significantly increased the cytosolic levels of GSH ($P < 0.05$) (Figure 1C). This result, together with the decreased ROS generation, reflects a diminished intracellular oxidation which could be expected to place the cell in favourable conditions to face a potential oxidative insult. As it is demonstrated in the present study, treatment of HepG2 cells with t-BOOH induced a significant decrease in the concentration of GSH ($P < 0.001$), as it was previously demonstrated (Goya, Mateos, & Bravo, 2007). However, this depletion of the GSH store was partly prevented by pre-treatment with lunasin (Figure 1C). These findings indicate that increased levels of GSH in the lunasin-treated cells before exposure to the oxidative damage greatly helped to prevent the dramatic depletion of intracellular GSH during the oxidative stress, an effect of lunasin that had not been reported previously. Maintaining GSH concentration above a critical threshold while facing a stressful situation represents an enormous advantage for cell survival.

### 3.2. Influence of lunasin on antioxidant enzymes activity

In order to investigate whether the antioxidant properties of lunasin are related to the induction of the antioxidant enzymes activity, HepG2 cells were treated with this peptide, and the activity of enzymes GPx and CAT was measured. In non-stressed HepG2 cells, lunasin did not provoke any change in the activity of GPx and CAT (data not shown). These results indicate that in spite of the direct effects of lunasin on ROS levels and GSH content, the defence enzymatic system in the cells was balanced. The increase in the activity of GPx and CAT observed after exposure to t-BOOH (Figure 3A and 3B), clearly indicates a positive response of the cell defense system to face the increasing generation of ROS evoked by the oxidative insult (Alía et al., 2006; Goya, Martín, Ramos, Mateos, & Bravo, 2009). It has been demonstrated that GPx and CAT play a crucial role as the first line of the antioxidant defense system against ROS generated during oxidative stress (Ray & Husain,
GPx catalyses reduction of $H_2O_2$ or other peroxides at the expense of GSH oxidation to oxidized glutathione, and CAT converts $H_2O_2$ to $H_2O$. Both enzyme activities are thus essential for the intracellular quenching of cell-damaging peroxide species but by two distinct mechanisms, since GPx is a glutathione related enzyme whereas CAT is independent from glutathione (Goya, Martín, Ramos, Mateos, & Bravo, 2009). However, a rapid return of the antioxidant enzyme activities to basal values once the challenge has been surmounted will place the cell in a favorable condition to deal with a new insult. Therefore, changes in their activity are considered as biomarkers of the antioxidant cellular response. In non-stressed HepG2 cells, lunasin did not provoke any change in the activity of GPx and CAT (data not shown). These results indicate that in spite of the direct effects of lunasin on ROS levels and GSH content, the defence enzymatic system in the cells was balanced. The increase in the activity of GPx and CAT observed after exposure to t-BOOH (Figure 3A and 3B), clearly indicates a positive response of the cell defense system to face the increasing generation of ROS evoked by the oxidative insult (Alía et al., 2006; Goya et al., 2009). However, a rapid return of the antioxidant enzyme activities to basal values once the challenge has been surmounted will place the cell in a favorable condition to deal with a new insult. It has been previously shown that flavonoid quercetin (Alía et al., 2006), olive oil phenol hydroxytyrosol (Goya et al., 2007), organic selenium derivatives (Cuello et al., 2007), and seaweed metabolite phloroglucinol (Quéguineur et al., 2012), among others, protect HepG2 cell integrity by preventing the severely increased activity of antioxidant enzymes induced by t-BOOH. In the present study, we show, for the first time, that a 20 h treatment of human HepG2 cells with lunasin prevents the permanent increase in the activity of both glutathione-related GPx and glutathione-independent CAT induced by oxidative stress (Figure 3A and 3B). Thus, the restrained ROS production during the stressful challenge in lunasin-treated HepG2 cells reduces the need of peroxide detoxification through GPx and CAT. Although a
potential direct effect of lunasin on antioxidant enzymes gene expression throughout the
antioxidant response element cannot be ruled out, the protective mechanism of lunasin can be
illustrated in terms of regulation of the specific activity of antioxidant defence enzymes.

3.3. Lunasin-induced reduction of protein oxidation

Since protein carbonyl concentration has been found elevated in various diseases thought
to be related to free radical damage, it has been widely used as an index of protein oxidation
in biological and medical sciences (Mateos, & Bravo, 2007). Figure 4A shows that 3 h-
treatment of HepG2 with 400 µM t-BOOH evoked a significant increase in the cellular
concentration of protein carbonyl groups of about 3-times compared to non-stressed cells,
indicating an intense oxidative damage to cell proteins. However, pre-incubation of the cells
with lunasin at concentrations ranged from 0.5 to 10 µM for 20 h reduced the protein carbonyl
levels down to levels measured in non-stressed cells. This fact indicates the ability of this
peptide to diminish the level of protein oxidation resulting from chemical induction with t-
BOOH. Other food compounds including plant polyphenols, beta carotene, lutein, seaweed
metabolite phloroglucinol and biscuit melanoidins have been also reported to prevent protein
oxidation (Alía et al., 2006; Goya et al., 2007; Martín, Ramos, Mateos, Izquierdo-Pulido,
Bravo, & Goya, 2010; Murakami, Hirakawa, Inui, Nakano, & Yoshida, 2002; Quéguineur et
al., 2012). However, to date, no dietary peptide had demonstrated to exert this protective
effect.

3.4. Lunasin prevents apoptosis by reducing stress-induced caspase-3

Oxidative stress-induced hepatic cell injury results not only from direct chemical
interactions by altering cellular macromolecules such as DNA, proteins and lipids, but also
from alterations in key mediators of stress signals and stress-dependent apoptosis reactions
In this regard, ROS generation has been described as a critical upstream activator of the caspase cascade that ends up with stimulation of downstream key effectors such as caspase-3 and subsequent development of apoptosis (Singh, & Czaja, 2008). Besides, GSH depletion is a common feature of apoptotic cells, and its role as a critical regulator in the signaling pathways leading to the progression of apoptosis has been reported (Franco, & Cidlowski, 2009, 2012). As shown in Figure 4B, the caspase-3 activity was significantly increased after 400 μM t-BOOH treatment for 3 h ($P < 0.001$), compared with the non-stressed cells. Previous studies had also demonstrated this pro-apoptotic effect of chemical t-BOOH on HepG2 cells through activation of caspase-3 and, ultimately, cell death (Martín, Granado-Serrano, Ramos, Izquierdo-Pulido, Bravo, & Goya, 2010). Consistent with the above mentioned ROS scavenging effect of lunasin, this peptide in the range of 1-10 μM effectively reduced caspase-3 activity in t-BOOH-induced HepG2 cells ($P < 0.001$) although no dose-response was observed, indicating that increases in lunasin’s concentration did not improved the caspase-3 activity reduction caused by this peptide. The demonstrated effects indicated the preventive capacity of lunasin against preventing the apoptotic effects induced by disruption of the redox steady-state.

### 3.5. Stability of peptide lunasin in HepG2 cultures

In order to study the stability of lunasin after its addition to HepG2 cells and incubation for 20 h, the medium without and with lunasin (10 μM) at different incubation hours (0, 2, 6, 12, and 20 h) was analyzed by HPLC-MS/MS. As an example, Figure 5 shows the extracted ion chromatograms of the molecular ion of lunasin $m/z$ 1257.5 (charge +4) obtained after 6, 12, and 20 h-incubation of cells with lunasin. The lunasin’s peak area was measured (Figure 6), and it was observed that the content of this peptide in the medium notably decreased with the incubation time. After 12 h-incubation, only 29% of initial lunasin remained in the
medium, and after 20 h, less than 1% was observable. In the present study, the analysis by HPLC-MS/MS also allowed the identification of lunasin-derived fragments that could be released during incubation of lunasin with HepG2 cells (Figure 5). Five lunasin-fragments were observed; all of them corresponding to the C-terminal region of the peptide, from the amino acid residue 25. The identified peptides’ peak areas were also measured (Figure 6). After 2 h-incubation, fragments f(25-43) and f(26-43) were already detectable in the medium. Other three peptides were released at 6 and 12 h-incubation. It has to be highlighted that after 20 h, extracellular lunasin only represented the 0.6% of total identified peptides whereas peptide f(25-43) represented 76.5%. Therefore, it can be postulated that the most abundant fragment, f(25-43), can be, at least, in part, responsible of the activity observed, and further studies with this lunasin-derived peptide are already in progress. In spite the important hydrolysis observed, it cannot be excluded that part of the lunasin could also internalize into HepG2 cells. It is also important to note that all the identified fragments contain the C-terminal part of the lunasin, which had previously been proposed as the active site of lunasin (Hernández-Ledesma et al., 2013). The hydrolysis of lunasin observed in contact with the HepG2 cells was different to that observed in our previous study with differentiated human intestinal Caco-2 cells (García-Nebot et al., 2014), where most of lunasin added to culture medium remained intact after 24 h incubation with these cells.

In the present study, analyses, focused on evaluating the antioxidant activity of peptide lunasin in HepG2 cells, were carried out after 20 h pre-treatment with this peptide. It was demonstrated the protective mechanism on cells submitted to an oxidative stress that can be illustrated in terms of regulation of the cellular redox status, i.e. peptide treatment restrained ROS production and maintained GSH concentration during the stress which reduced the necessity of peroxide detoxification through GPx and CAT. Additionally, a controlled ROS generation reduced oxidative damage to proteins and restrained activity of the apoptotic
pathway resulting in improved cell viability. The final findings found on lunasin’s stability in HepG2 cells indicated that this peptide was markedly hydrolyzed at the selected time (20 h), and thus, in addition to the remaining peptide, the fragments released during its hydrolysis might be the actual responsible for the observed effects. All the peptides identified at 20 h corresponded to the active site-sequence of lunasin described to date. Also, they contained the motif RGD known to be crucial for the interaction of proteins or peptides with its cell surface receptor (Ruoslahti, & Pierschbacher, 1986). Therefore, these structural characteristics make these five peptides promising chemo-protective peptides against oxidative stress in liver HepG2 cells. In conclusion, our results point out for the first time a direct antioxidant action of lunasin or its derived fragments on hepatocytes exposed to oxidizing species. It indicates the possibility that these peptides may significantly contribute to preserve the integrity of liver tissues against oxidative damage related disorders. Since this study has been carried out with synthetic lunasin which secondary and tertiary structure could differ to that of plant-purified lunasin, confirmation of the effects should be needed to demonstrate the chemo-protective potential of natural lunasin.

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Conflict of interests

The authors declare no competing financial interest.
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FIGURE CAPTIONS

Figure 1. Direct and protective effects of peptide lunasin on cell viability and intracellular ROS and GSH levels. HepG2 cells were pre-incubated with medium or medium supplemented with lunasin (final concentration ranged from 0.5 to 10 μM) for 20 h before treatment with medium (□: non-stressed cells) or medium supplemented with 400 μM t-BOOH for 3 h (■). (A) Cell viability was measured by the CV assay. Results are expressed as percent of viable cells, n = 12. (B) Intracellular ROS generation. Fluorescence units corresponding to intracellular ROS production are expressed as percent of data from non-stressed cells, n = 8. (C) Intracellular GSH levels. Results of fluorescent analysis are calculated as nmoles of GSH per mg of protein, and represented as percentage of non-stressed cells, n = 6. * (P < 0.05), *** (P < 0.001), significantly different from control non-stressed cells, and ### (P < 0.001), significantly different from control t-BOOH-induced cells.

Figure 2. Morphological analysis of HepG2 cells. HepG2 cells were pre-incubated with medium or medium supplemented with lunasin (final concentration 0.5-10 μM) for 20 h before treatment with medium (non-stressed cells) or medium supplemented with 400 μM t-BOOH for 3 h. Representative images of (A) non-stressed cells pre-incubated with medium, (B) t-BOOH-induced cells pre-incubated with medium, and t-BOOH-induced cells pre-incubated with (C) 0.5 μM lunasin and (D) 5 μM lunasin. HepG2 cells pre-incubated with medium and treated with t-BOOH showed reduction in cell numbers and loss of cell-to-cell contact (arrows), compared with non-stressed cells pre-incubated with medium, and t-BOOH-induced cells pre-incubated with lunasin. Size bar: 10 μm.

Figure 3. Protective effects of lunasin on enzymes GPx and CAT activity. HepG2 cells were pre-incubated with medium or medium supplemented with lunasin (final concentration ranged...
from 0.5 to 10 μM) for 20 h before oxidation induction with 400 μM t-BOOH for 3 h (■). Results of enzymes (A) GPx and (B) CAT activities are calculated as mUnits per mg of protein and represented as percentage of data from non-stressed cells (□). Represented values are means ± SD of n = 6 (GPx) and n = 4 (CAT). *** (P < 0.001), significantly different from control non-stressed cells, and # (P < 0.05), ## (P < 0.01), ### (P < 0.001), significantly different from control t-BOOH-induced cells.

Figure 4. Protective effects of lunasin on protein carbonyl content and caspase-3 activity. HepG2 cells were pre-incubated with medium or medium supplemented with lunasin (final concentration ranged from 0.5 to 10 μM) for 20 h before oxidation induction with 400 μM t-BOOH for 3 h (■). (A) Protein carbonyl content. Results are expressed as nmol per mg protein and represented as percentage of non-stressed cells (□). (B) Caspase-3 activity. Results are calculated as Units per μg of protein and expressed as percent of control data. Values are means ± SD of 4 different samples per condition. *** (P < 0.001), significantly different from control non-stressed cells, and ### (P < 0.001), significantly different from control t-BOOH-induced cells.

Figure 5. Stability of peptide lunasin in medium added to HepG2 cells. (A) Extracted ion chromatogram (EIC) of the molecular ion of lunasin (F1) m/z 1257.5 (charge +4), f(32-43) (F2) m/z 1324.5 (charge +1), f(30-43) (F3) m/z 1565.5 (charge +1), f(29-43) (F4) m/z 1693.8 (charge +1), f(26-43) (F5) m/z 1034.1 (charge +2), f(25-43) (F6) m/z 1102.7 (charge +2) in serum free DMEM-F12 medium incubated with 10 μM lunasin and collected after (A) 6 h, (B) 12 h, and (C) 20 h-incubation. (B) Relative amount (expressed as peak area) of lunasin (F1) and its derived fragments (F2-F6) in serum free DMEM-F12 medium incubated with lunasin for 0, 2, 6, 12, and 20 h.