Cocoa flavonoids improve insulin signalling and modulate glucose production via AKT and AMPK in HepG2 cells

Isabel Cordero-Herrera¹, María Angeles Martín¹², Laura Bravo¹, Luis Goya¹ and Sonia Ramos¹,*

¹Department of Metabolism and Nutrition
Institute of Food Science and Technology and Nutrition (ICTAN)
Consejo Superior de Investigaciones Científicas (CSIC)
José Antonio Novais 10
Ciudad Universitaria, 28040, Madrid
Spain
Phone: +34.91.544.56.07
Fax: +34.91.549.36.27
²Centro de Investigación Biomédica en Red de Diabetes y Enfermedades Metabólicas Asociadas (CIBERDEM), ISCIII, Spain.

* Corresponding author: e-mail: s.ramos@ictan.csic.es

Abbreviations used: AKT/PKB, protein kinase B; AMPK, 5´-AMP-activated protein kinase; BrdU, 5-bromo-2´-deoxyuridine; CaMKK, Ca²⁺/calmodulin-dependent protein kinase kinase; CPE, cocoa phenolic extract; EC, (-)-epicatechin; EGCG, epigallocatechin gallate; FBS, fetal bovine serum; FOXO1, forkhead box protein O1; GLUT, glucose transporter; GS, glycogen synthase; GSK-3, glycogen synthase kinase-3; G6Pase, glucose-6-phosphatase; IR, insulin receptor; IRS, insulin receptor substrate; LKB1, liver kinase B1; PPAR, peroxisome proliferator-activated receptor; PEPCK, phosphoenolpyruvate carboxykinase; PI3K, phosphatidylinositol-3-kinase.

Keywords: Cocoa, Epicatechin, Glucose production, Insulin signalling pathway, HepG2 cells.
Abstract

Scope: Cocoa and (-)-epicatechin (EC), a main cocoa flavanol, have been suggested to exert beneficial effects in diabetes, but the mechanism for their insulin-like effects remains unknown. In this study, the modulation of insulin signalling by EC and a cocoa phenolic extract (CPE) on hepatic HepG2 cells was investigated by analysing key proteins of the insulin pathways, namely insulin receptor (IR), insulin receptor substrate (IRS)-1 and 2, PI3K/AKT and 5´-AMP-activated protein kinase (AMPK), as well as the levels of the glucose transporter GLUT-2 and the hepatic glucose production.

Methods and results: EC and CPE enhanced the tyrosine phosphorylation and total IR, IRS-1 and IRS-2 levels and activated the PI3K/AKT pathway and AMPK in HepG2 cells. CPE also enhanced the levels of GLUT-2. Interestingly, EC and CPE modulated the expression of phosphoenolpyruvate carboxykinase, a key protein involved in the gluconeogenesis, leading to a diminished glucose production. In addition, EC- and CPE-regulated hepatic gluconeogenesis was prevented by the blockage of AKT and AMPK.

Conclusion: Our data suggest that EC and CPE strengthen the insulin signalling by activating key proteins of that pathway and regulating glucose production through AKT and AMPK modulation in HepG2 cells.
1. Introduction

Diabetes is one of the most common chronic diseases in nearly all countries, and it is continuing to be an increasing international health burden [1]. Current medications are not adequately effective in maintaining long-term glycemic control in most patients. Therefore, there is an urgent need to continue working on the prevention and control of diabetes. In this regard, a promising approach is the employ of the natural compounds with insulin-like activity, which have been proposed as potential therapeutic agents in the prevention and/or treatment of this disease [2].

The liver has an important role in the control of the whole body metabolism of energy nutrients. In diabetes, the insulin target tissues are damaged, which aggravates the ability of insulin to trigger downstream metabolic actions resulting in insulin resistance. In this regard, one of the hallmarks of diabetes is the alteration of the hepatic metabolism: the liver is not able to control the glucose homeostasis, which is one of the causes for the hyperglycemia, and there is a miss-regulation of the insulin pathway in this organ [3].

Insulin signal transduction is initiated when insulin binds to the insulin receptor (IR), which leads to the stimulation of several intracellular protein substrates including insulin receptor substrate (IRS)-1 and IRS-2. This triggers the phosphatidylinositol-3-kinase (PI3K) pathway that stimulates AKT and leads to the inhibition of glycogen synthase kinase-3 (GSK-3) by phosphorylation, which subsequently phosphorylates and inactivates glycogen synthase (GS), and conducts to the modulation of other proteins necessary for the acute metabolic effects of insulin [3]. In addition, in the hepatocyte AKT might phosphorylate the transcription factor forkhead box protein O1 (FOXO1), restraining the expression of phosphoenolpyruvate carboxykinase (PEPCK) and glucose-6-phosphatase (G6Pase), and inhibiting the gluconeogenesis [3]. 5’-AMP-
activated protein kinase (AMPK) also constitutes a central regulator of the cellular metabolism, able to suppress the hepatic gluconeogenesis [4]; in fact numerous drugs that are in clinical use for treatment of type 2 diabetes activate AMPK [4, 5].

(-)-Epicatechin (EC) is one of the most abundant flavonoids in human diet, being present in high concentrations in grapes, cocoa, tea, and many other fruit and vegetables [6, 7]. Different studies have shown that EC is able to interfere with the oxidative/antioxidative potential of hepatic cells [6, 8], induces survival/proliferation pathways [8-10], exerts insulin-like activities [11], and improves insulin sensitivity [12], blood glucose levels [13] and parameters related to the inflammation in cardiovascular disease and diabetes [12, 14, 15]. Similarly, dark chocolate or cocoa, which are widely consumed [16] and are a rich source of flavonoids, especially (-)-epicatechin [6], have been reported to protect against carcinogenesis [7, 17], improve the allergy process [18], cardiovascular status [16, 19, 20], insulin sensitivity and hyperglycemia [19, 21] in humans and experimental animals. However, the precise mechanism for the preventive activities of EC and cocoa related to glucose metabolism and insulin signalling in the liver remains largely unknown.

The present work evaluates the effects of EC and a cocoa phenolic extract (CPE) on insulin signalling pathways and glucose production in HepG2 cells. The study demonstrates that EC and CPE have insulinomimetic activities in the human hepatic HepG2 cells, as EC and CPE enhanced the tyrosine phosphorylation and total IR, IRS-1 and IRS-2 levels, which was associated to an activation of the PI3K/AKT pathway and AMPK. We have also found that AKT and AMPK are required to modulate the hepatic glucose production.
2. Materials and methods

2.1. Materials and chemicals

(-)-EC (>95% of purity), Compound C (6-[4-(2-Piperidin-1-ylethoxy)phenyl]-3-pyridin-4-ylpyrazolo[1,5-a]pyrimidine), LY294002 (2-(4-morpholinyl)-8-phenyl-1(4H)-benzopyran-4-one hydrochloride), anti-mouse IgG-agarose, sodium lactate, sodium pyruvate, gentamicin, penicillin G and streptomycin were purchased from Sigma Chemical (Madrid, Spain). Cell proliferation ELISA BrdU (colorimetric) assay kit was from Roche Diagnostics (Barcelona, Spain). Anti-AKT and anti-phospho-Ser473-AKT detecting levels of phospho- and total AKT 1-3, anti-AMPK and anti-phospho-Thr172-AMPK, as well as anti-GSK3 α/β and anti-phospho-GSK3 α/β recognizing phosphorylated Ser21/9 of GSK3, anti-GS and anti-phospho-GS recognizing phosphorylated Ser641 of GS, anti-IRS-2 and anti-β-actin were obtained from Cell Signalling Technology (Izasa, Madrid, Spain). Anti-IR β, anti-PEPCK and anti-Tyr(P) (PY20) were purchased from Santa Cruz (sc-711, sc-32879 and sc-508, respectively, Qimigen, Madrid, Spain). Anti-IRS-1 and anti-GLUT-2 were from Millipore (Madrid, Spain). Materials and chemicals for electrophoresis were from BioRad (BioRad Laboratories S.A., Madrid, Spain). Cell culture dishes and cell culture medium were from Falcon (Cajal, Madrid, Spain) and Lonza (Madrid, Spain), respectively.

2.2. Cocoa polyphenol extraction

Natural Forastero cocoa powder (Nutrexpa, Barcelona, Spain) was used for this study. Soluble polyphenols were extracted by sequentially washing 1 g of sample with 40 mL of 16 mM hydrochloric acid in 50% aqueous methanol (50:50, v/v, 1 h at room temperature, constant shaking) and 40 mL of acetone:water (70:30, v/v, 1 h at room temperature, constant shaking). After centrifugation (15 min, 3000 g), supernatants
from each extraction step were combined and made up to 100 mL. The desiccated extract was dissolved in distilled water and kept frozen until assay. A detailed description of this cocoa polyphenol extract (CPE) is given elsewhere [6, 22]. Accordingly, the amount of EC and polyphenols present in the CPE were 383.5 mg/100 g (determined by LC-MS) and 2 g/100 g on dry matter basis (determined by Folin-Ciocalteu) [6]. Concentrations of EC in tested doses of CPE range from 13.2 nM (in the dose of 1 µg CPE/mL) to 132 nM (in that of 10 µg CPE/mL) [6].

2.3 Cell culture and treatments

Human HepG2 cells were grown in DMEM-F12 medium supplemented with 2.5% foetal bovine serum (FBS) and the following antibiotics: gentamicin, penicillin and streptomycin (50 mg/L). Cells were maintained at 37°C in a humidified atmosphere of 5% CO₂. One day after plating, the medium was changed to DMEM containing 5.5 mM D-glucose, 2 mM glutamine and FBS, and the culture was continued. Subsequently, the experimental treatment was carried out for the indicated periods with various concentrations of EC or CPE in serum-free media containing 5.5 mM D-glucose, 2 mM glutamine for 24 h.

Cells were treated with different concentrations of EC (1-20 µM) or CPE (1-20 µg/mL) diluted in serum-free culture medium during 24 h. In the experiments with the pharmacological inhibitors, cells were preincubated with 25 µM LY294002 or 40 µM Compound C for 1 h prior to 24 h of EC or CPE treatment.

2.4. Cell viability assay

Cell viability was determined by using the crystal violet assay [8]. HepG2 cells were seeded at low density (10⁴ cells per well counted in a Neubauer chamber) in 96-well
plates, grown for 20 h and incubated with crystal violet (0.2% in ethanol) for 20 min. Plates were rinsed with water, allowed to dry, and 1% sodium dodecylsulfate added. The absorbance of each well was measured using a microplate reader at 570 nm.

2.5. Cell proliferation assay (5-bromo-2’-deoxyuridine assay, BrdU)

A colorimetric immunoassay (ELISA) was used for the quantification of cell proliferation [8]. This method is based on the measurement of BrdU incorporation into genomic DNA during DNA synthesis of proliferating cells. HepG2 cells were seeded (10^4 cells per well counted in a Neubauer chamber) in 96-well plates, grown 20 h and labelled by the addition of BrdU for 4 h. Then the anti-BrdU antibody was added and the immune complexes were quantified by measuring the absorbance at 620 nm in a microplate reader.

2.6. Preparation of cell lysates

Cells were lysed at 4°C in a buffer containing 25 mM HEPES (pH 7.5), 0.3 M NaCl, 1.5 mM MgCl₂, 0.2 mM EDTA, 0.5 mM dithiothreitol, 0.1% Triton X-100, 200 mM β-glycerophosphate, 0.1 mM Na₃VO₄, 2 µg/mL leupeptin and 1 mM phenylmethylsulfonyl fluoride. The supernatants were collected, assayed for protein concentration by using the Bio-Rad (Bio-Rad, Madrid, Spain) protein assay kit according to the manufacture’s specifications, aliquoted and stored at -80°C until used for immunoprecipitation and/or Western blot analyses.

2.7. Immunoprecipitation

Protein extracts containing 200 µg of protein were immunoprecipitated overnight at 4°C with gentle rotation in the presence of 2-5 µg of anti-Tyr(P) (PY20) antibody, followed
by the addition of anti-mouse IgG-agarose. After mixing for 2 h, the pellets were collected by centrifugation, and the supernatants were discarded. Then the pellets were washed and saved for Western blot analyses.

2.8. Western blot analysis

Equal amounts of proteins were separated by SDS-polyacrylamide gel electrophoresis and transferred to polyvinylidene difluoride filters (Bio-Rad, Madrid, Spain). Membranes were probed with the corresponding primary antibody followed by incubation with peroxide-conjugated anti-rabbit (GE Healthcare, Madrid, Spain) or anti-mouse (Sigma, Madrid, Spain) immunoglobulin. Blots were developed with the ECL system (GE Healthcare, Madrid, Spain). Normalization of Western blot was ensured by β-actin and bands were quantified using a scanner and accompanying software.

2.9. Glucose production assay

HepG2 cells were seeded in 24-well plates (2 x 10^5 cells per well counted in a Neubauer chamber) and treated in serum-free DMEM with 10 µM EC or 1 µg/mL CPE for 24 h. The medium was then replaced with glucose production buffer consisting of glucose-free DMEM (pH 7.4), without phenol red (Invitrogen, Madrid, Spain), supplemented with 20 mM sodium lactate and 2 mM sodium pyruvate, as previously described [23, 24]. After a 3-h incubation, medium was collected and glucose concentration measured with a colorimetric glucose assay kit (Sigma, Madrid, Spain). The readings were then normalized to the total protein content determined from the whole-cell lysates.

2.10. Statistics
Prior to statistical analysis, data were tested for homogeneity of variances by the test of Levene; for multiple comparisons, one-way ANOVA was followed by the Bonferroni test when variances were homogeneous or by the Tamhane test when variances were not homogeneous. $P<0.05$ was considered significant. A SPSS version 19.0 program has been used.
3. Results

3.1. Cell viability and proliferation

To determine the potential effects on cell viability and proliferation of EC and CPE in a human hepatic cell line (HepG2), cells were exposed to a range of concentrations (0-20 µM and 0-20 µg/mL, respectively) for 24 h.

Treatment of HepG2 cells for 24 h with EC or CPE did not evoke changes in cell viability, as determined by the crystal violet assay, indicating that the concentrations selected for the study did not damage cell integrity during the period of incubation (Table 1). Similarly, treatment with EC or CPE did not affect cell growth, indicating no impairment of cell proliferative machinery and preservation of a regular cell cycle (Table 1). Since none of the tested doses induced cell toxicity, the lowest and realistic range of concentrations was selected for further studies [6, 22].

3.2. Effects of EC and CPE on tyrosine phosphorylation and protein levels of IR and its substrates IRS-1 and IRS-2

The modulation of IR and its downstream substrates IRS-1 and IRS-2 is essential for recruiting and activating downstream pathways [25]. In fact, in hepatic insulin resistance the insulin-stimulated-IR and IRS tyrosine phosphorylation is defective and results in reduced IRS-associated PI3K activities [3]. To test the effect of EC and CPE on tyrosine phosphorylation and total levels of IR and its substrates, HepG2 cells were exposed for 24 h to various concentrations of these substances.

As shown in Figure 1, IR and IRS-2 phosphorylated and total protein levels equally increased with all concentrations of EC and CPE tested, as well as the phosphorylated and total protein levels of IRS-1 when cells were incubated with CPE. EC dose-dependently increased the phosphorylated and total levels of IRS-1 (Figures 1A and
1C). All these results suggest that EC and CPE could strengthen the insulin signalling cascade by up-regulating the early steps of this pathway.

3.3. Effects of EC and CPE on AKT, GSK3 and GS phosphorylation

AKT is the molecular key in mediating the metabolic effects of insulin signalling [25]. It lays downstream of PI3K and facilitates glucose uptake and glycogen synthesis in the liver [25]. To test the modulation of AKT by EC and CPE, phosphorylated and total AKT were evaluated in cell lysates by Western blot analysis. As shown in Figures 2A and 2B, the treatment of cells with EC and CPE for 24 h increased the phosphorylation of AKT with all concentrations tested. EC (10 µM) showed higher levels of p-AKT than lower concentrations of EC, whereas all doses of CPE induced a similar enhancement in p-AKT values.

Since AKT directly contributes to the activity of GS [25], the effect of EC and CPE on the phosphorylated and total protein expression levels of GSK3 and GS was assayed. Western blot analysis of GSK3 and p-GSK3 proteins showed two bands corresponding to the α and β isoforms. EC and CPE treatment enhanced the phosphorylated isoforms, as illustrated in Figures 2A and 2C. EC (10 µM) induced the highest increase in the levels of p-GSK3 in comparison to the other doses of EC, and CPE equally enhanced the levels of p-GSK3 with all concentrations assayed, in agreement to what was observed for AKT (Figures 2A and 2B).

As shown in Figures 2A and 2D, p-GS levels decreased in the presence of EC or CPE with all concentrations tested. As it could be expected according to the previous results, CPE equally diminished the phosphorylated levels of GS and 10 µM EC induced the most remarkable diminution of p-GS of all EC concentrations assayed. There was no difference in the total levels of AKT, GSK3 and GS.
3.4. Effects of EC and CPE on AMPK phosphorylation

To continue the study of the potential effect of EC and CPE on key proteins of the insulin signalling, AMPK, which is a sensor of energy status for maintaining cellular energy homeostasis [5], was evaluated by analysing the total and phosphorylated levels in total cell lysates by Western blot analysis. Figure 3 illustrates that a 24 h treatment with EC or CPE induced an increase in AMPK phosphorylated levels. EC (5 and 10 µM) showed higher levels of p-AMPK than the lowest concentration of EC, whereas CPE induced a similar activation of AMPK with all concentrations tested. The protein levels of total AMPK were not modified by EC or CPE treatment.

3.5. Effects of EC and CPE on GLUT-2 protein levels

The GLUT-2 transporter mediates the diffusion of glucose across the plasma membrane of the hepatocyte, maintaining intracellular glucose in equilibrium with extracellular glucose [3]. To test whether this transporter was affected by EC and CPE, HepG2 cells were incubated with the selected concentrations for 24 h. Treatment of HepG2 cells with any of the three concentrations of EC did not affect the expression levels of GLUT-2, and only 10 µg/mL CPE increased the levels of GLUT-2 after a 24 h-incubation (Figure 4).

3.6. Effect of EC and CPE activation of AKT on hepatic gluconeogenesis

In the liver, insulin-activated AKT inhibits the expression of PEPCK and G6Pase and, therefore, gluconeogenesis [3]. In view of the increased levels of p-AKT induced by EC and CPE, it was studied whether both substances were able to modulate the expression of a major enzyme responsible of the regulation of gluconeogenesis, such as PEPCK, as
well as the production of glucose. To this end, HepG2 cells were exposed to a selective inhibitor of AKT (LY294002) and EC or CPE and the levels of PEPCK and the novo production of glucose were assayed. The concentrations selected for these analyses were the doses that exhibited a prominent effect on the activation of the PI3K/AKT pathway, i.e.: 10 µM for EC and 1 µg/mL for CPE.

Treatment of HepG2 cells with EC or CPE showed a comparable decrease in the expression of PEPCK (Figures 5A and 5B). LY294002 alone increased the levels of PEPCK, although these values decreased by the addition of EC and CPE (Figures 5A and 5B). LY294002 treatment decreased p-AKT levels in all cells incubated with this inhibitor, although p-AKT values were partly recovered in the presence of EC or CPE when compared to untreated cells (Figures 5A and 5C). p-AMPK and total expression of AKT and AMPK were not affected by LY294002 (Figures 5A, 5C and 5D). EC and CPE decreased the production of glucose (Figure 5E). In addition, the presence of the AKT selective inhibitor alone increased the hepatic glucose production, and this enhancement was less prominent in LY+EC- and LY+CPE-treated cells (Figure 5E). Conversely, LY294002 could not totally block the inhibitory effect of EC and CPE on PEPCK and glucose production, indicating that other pathways may also be involved in EC- and CPE-induced effects.

3.7. Effect of EC and CPE activation of AMPK on hepatic gluconeogenesis

AMPK is another known suppressor of hepatic gluconeogenesis [3]. To examine the role of AMPK on the modulation of the levels of PEPCK and the production of glucose, AMPK was blocked by a selective chemical inhibitor (Compound C) prior to the treatment with EC or CPE. As mentioned above, the concentrations selected for these
analyses were the previously chosen (10 µM for EC and 1 µg/mL for CPE), since they showed a prominent effect on the activation of AMPK.

As illustrated in Figures 6A and 6B, EC and CPE decreased the levels of PEPCK, and the blockage of AMPK induced a significant increase in the expression of the mentioned gluconeogenic enzyme. However, this enhancement was less pronounced in the presence of EC and CPE in comparison to that induced by the compound C alone. Treatment with compound C alone decreased p-AMPK levels, whereas incubation with EC or CPE and compound C partly recovered p-AMPK levels (Figures 6A and 6D). p-AKT and total AKT and AMPK remained unaltered after incubating the cells with compound C (Figures 6A, 6C and 6D). In addition, compound C enhanced the glucose production via hepatic gluconeogenesis (Figure 6E). Similarly, the increase in glucose production was higher in compound C-treated cells than in C+EC- and C+CPE-treated cells (Figure 6E). However, compound C could not completely block the inhibitory effect of EC and CPE on PEPCK and glucose production, indicating that other pathways may also contribute to the beneficial effects of EC and CPE.
4. Discussion

Flavonoids have been found to possess beneficial effects on health and have drawn attention because of their safety and accumulating evidence on their antidiabetic effects in animals and humans [2]. In the current study, we have demonstrated for the first time that EC and CPE enhanced the tyrosine phosphorylated and total IR, IRS-1 and IRS-2 levels together with an activation of the PI3K/AKT pathway and AMPK (Figure 7). We have also provided evidence on the insulin-like activity of EC and CPE, which were able to down-regulate the levels of the key gluconeogenic enzyme PEPCK and modulate the hepatic glucose production through AKT and AMPK (Figure 7). HepG2 cells are widely used for biochemical and nutritional studies as a cell culture model of human hepatocytes since they retain their morphology and most of their function in culture [26-28]. Thus, this cell line has been extensively used to study the hepatic glucose production and the modulation of the insulin pathway in vitro [27-30].

The liver plays a critical role in maintaining blood glucose concentration both through its ability to supply glucose to the circulation via glycogenolysis and gluconeogenesis in the postabsorptive state and to remove glucose from the circulation after meal ingestion [3]. However, in diabetes the gluconeogenic pathway is aberrantly activated, supplies a relatively larger amount of glucose into the circulation [2, 3], and there is also hepatic insulin resistance [3]. Therefore, the modulation of the mentioned targets could be beneficial for the prevention and control of this disease; in this line, natural compounds such as flavonoids could play a major role, although little investigation at the molecular level has been performed.

In the present study it is shown that EC and CPE increased tyrosine phosphorylation and total levels of IR and IRS in HepG2 cells. In concert, epigallocatechin gallate (EGCG) and naringenin attenuated high glucose-induced signalling blockage by
reducing IRS-1 serine phosphorylation in HepG2 cells [27] and by activating IRS-1 in primary hepatocytes of mice with metabolic syndrome [31]. In addition, green tea polyphenols increased IRS-2 mRNA levels in the liver [32], as well as IR and IRS-1 and -2 levels in the myocardium of insulin-resistant rats [33]. Similarly, oligomers of a grape-seed procyanidin extract increased IR and IRS levels in preadipocytes [34]. In this line, the oligomeric structures of the extract activated the IR by interacting with and inducing its tyrosine phosphorylation [34]. ECGC and rutin stimulated the IRS2 signalling by enhancing tyrosine phosphorylation under high glucose condition on pancreatic β cells [35], and in 3T3-L1 preadipocytes a CPE did not affect the levels of IR but it modulated its IR kinase activity via direct binding [36].

The activation of IRS-1/-2 initiates the stimulation of the PI3K/AKT pathway, which is needed for the metabolic effects of insulin in the liver and is responsible of the inhibition of GSK-3 and activation of GS [3]. In this regard, EC and CPE seemed to mimic the metabolic actions of insulin, as they increased the phosphorylated levels of AKT and GSK-3, and decreased p-GS values. Previous studies have also demonstrated the stimulation of AKT by EC and CPE in HepG2 cells [10, 22], and EC was reported to partly reverse the inhibition of AKT phosphorylation induced by high glucose levels [27]. Similarly, naringenin activated the AKT pathway in HepG2 cells [29] and green tea polyphenols increased the RNA expression levels of PI3K/AKT in the liver of insulin-resistant rats [32]. In this line, green tea polyphenols also increased the mRNA levels of GSK3 in the liver of insulin-resistant rats [32]. However, a decrease in the expression of GSK3 and an increase in the mRNA levels of GS have been reported in the myocardium of insulin-resistant rats [33]. All together indicates that polyphenols reinforced the signal pathways responsive to insulin.
In the liver, GLUT-2 is a glucose-sensitive gene that mediates both influx and efflux of glucose across the plasma membrane [3]. In the present study, levels of GLUT-2 were induced by the highest concentration of CPE tested, whereas EC did not show any effect on the modulation of the protein levels of this transporter. Although the influence of the phenolic compounds on the GLUTs has been scarcely evaluated, especially in the liver, in agreement with the present results no effects on the mRNA GLUT-2 levels in the liver of insulin-resistant rats fed with green tea polyphenols have been reported [32]. However, it should be mentioned that, as occurs for 10 µg/mL CPE, the grape polyphenol resveratrol and the antidiabetic drugs pioglitazone and rosiglitazone, which are peroxisome proliferator-activated receptor (PPAR)-γ agonists, were able to increase GLUT-2 expression in β-cells [37, 38]. In this line, our results could suggest that CPE could more efficiently contribute to glucose uptake from the blood when glucose levels are postprandially elevated in comparison with EC, as described for rosiglitazone [37]. This effect has been related to the regulation of PPARγ for rosiglitazone [37], as well as to the activation of AKT or AMPK in diabetic animals [39, 40], but further studies are needed to elucidate how GLUT-2 is modulated by CPE in hepatic cells.

AMPK is an important therapeutic target for diabetes as it is one of the central regulators of cellular metabolism, which can be activated at least by two pathways: liver kinase B1 (LKB1) and Ca^{2+}/calmodulin-dependent protein kinase kinase (CaMMK) [5]. In the present work, the phosphorylated levels of AMPK increased after incubating HepG2 cells with EC and CPE. Interestingly, the major green tea compound EGCG has previously been shown to activate AMPK in hepatic cells via CaMMK [23, 27]. In addition, other polyphenols such as theaflavins, naringin, anthocyanins, resveratrol and apigenin have proved to activate AMPK in the liver of mice and rats and, consequently, to modulate cellular metabolism [30, 31, 41, 42]. Similarly, other natural compounds
such as berberine, ginsenosides obtained from ginseng and extracts from the plant *Artemisia sacrorum* have been demonstrated to stimulate AMPK in HepG2 cells [43-46].

PEPCK is one of the major enzymes responsible of the regulation of gluconeogenesis [3]. In the present work, EC and CPE decreased PEPCK levels in HepG2 cells. In agreement with our results, other phenolic compounds such as EGCG, naringenin, catechin-rich green tea, naringin, genistein, daidzein and anthocyanins diminished the levels of this gluconeogenic enzyme in hepatic cells and mouse liver during an induced insulin-resistant situation and, consequently, reduced glucose production [23, 31, 41, 47-50].

AKT and AMPK are known to suppress the gluconeogenesis in the liver [3, 4, 25]. Suppression of hepatic gluconeogenesis by AKT-dependent insulin signalling is reduced or lost in type 2 diabetes due to insulin resistance plus a relatively insufficient insulin production [25]. In addition, activation of AMPK results in enhanced fatty acid oxidation and decreased production of glucose, cholesterol and triglycerides in the liver [3, 4]. Thus, activation of AMPK supresses G6Pase and PEPCK and then decreases hepatic glucose production [3, 4]. In agreement, we have shown that both kinases are involved in the modulation of PEPCK levels and the production of glucose in HepG2 cells treated with EC and CPE (Figure 7). As previously mentioned, EGCG activated AMPK and consequently inhibited hepatic gluconeogenesis via AMPK, although AKT seemed not to be involved in this process since it was not stimulated by the flavanol [23]. Similarly, naringenin did not modify AKT phosphorylation, and the incubation of the cells with LY294002 did not suppress the hepatic glucose production [48]. In addition, AMPK also mediated naringin repression of hepatic gluconeogenesis [31]. In this line, other natural compounds such as berberine, ginesenosides and an extract from
*Artemisia sacrorum* suppressed the hepatic glucose production and/or downregulated the levels of PEPCK via AMPK, although their effects on the mentioned parameters in the presence compound C were weaker than the effect showed by the natural compound alone [43-46]. Interestingly, clove extract acted like insulin in HepG2 cells by reducing PEPCK gene expression and this feature was reverted by LY294002 [51]. It has also been shown that LY294002 inhibited the expression of PEPCK, but AKT did not modulate the hepatic production of glucose via gluconeogenesis [23, 49]. Likewise, in the present study p-AMPK and p-GSK3β levels were increased by EC and CPE incubation. Accordingly, it has been showed that in the liver the activated form of AMPK is responsible for metabolic changes via phosphorylation of downstream substrates such as GSK3β and cAMP response element binding protein, which are directly or indirectly related to glucose production [43, 46]. In this regard, EGCG promotes phosphorylation of CaMKK, and blockade of CaMKK activity prevents EGCG activation of AMPK and mitigates the inhibitory role of EGCG in hepatic gluconeogenesis [23].

As mentioned above, cocoa is a rich source of flavonoids such as (-)-EC, (+)-catechin, and procyanidins, and EC is the most abundant flavanol in the CPE employed in this study [6]. Considering that concentrations of EC in tested doses of CPE range from 13.2 nM (in the dose of 1 µg CPE/mL) to 132 nM (in that of 10 µg CPE/mL) [6], and CPE effects are equivalent to those of the pure flavanol, it is reasonable to assume that EC is just one of the many bioactive substances present in CPE and that the synergic effect of phenolic compounds in foodstuffs should be taken into account, as previously shown [52].

It is worth mentioning that the range of concentrations used in the study is not far from realistic. In this regard, rats fed with EC showed plasma concentrations of EC and EC
metabolites of about 35 µM 1 h after oral administration of 172 µmol EC/Kg body weight [53]. In humans, levels of 6 µM EC and 41 nM procyanidin B2 have been reported after ingestion of 26 g cocoa [54]. Similarly, levels of 0.2–0.4 µM EC have been observed after ingestion of 50 g [55] and 80 g [56] chocolate. In addition, it should not be underestimated the potential contribution of EC and flavanol metabolites to the biological activity, which is unclear at present; their evaluation will require further studies. In this regard, more accurate approaches recently used are the incubation of cultured cells with plasma obtained from volunteers consuming the molecule or food of interest [57] and the employ of a system based on co-culture of human enterocytes with human hepatocytes, which has been proved to resemble a human physiological system useful for assay the bioactivity of extracts [58].

In summary, EC and CPE possess an insulin-like activity, as they enhanced tyrosine phosphorylated and total levels of IR, IRS-1 and IRS-2 and activated PI3K/AKT pathway and AMPK at concentrations that are not toxic to hepatic cells and are reachable through the diet (Figure 7). We have also revealed a new mechanism by which EC and CPE modulate the hepatic gluconeogenesis and PEPCK expression via AKT and AMPK (Figure 7). Although further efforts are needed to define the precise role of EC and cocoa in the regulation of the insulin pathways in liver cells, a diet rich in EC and/or cocoa may be a potential chemopreventive tool useful for the management of diabetes.
Acknowledgements

This work was supported by the grants AGL2010-17579 and CSD2007-00063 from the Spanish Ministry of Science and Innovation (MICINN). I. Cordero-Herrera is a fellow of the FPI predoctoral program of MICINN.

Conflict of interest statement

The authors have declared no conflict of interest.
5. References


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Legends to figures

**Figure 1.** Effect of EC and CPE on phosphorylated and total levels of IR, IRS-1 and IRS-2 in HepG2 cells after 24 h of treatment. (A) Bands of representative experiments. Densitometric quantification of (B) p-IR and total IR, (C) p-IRS-1 and total IRS-1 and (D) p-IRS-2 and total IRS-2. Protein extracts were subjected to immunoprecipitation (IP) with the anti-phospho-tyrosine (P-Tyr) antibody. The resulting immunocomplexes were analysed by Western blot (WB) with the anti-IR or IRS-1 or IRS-2 antibody. Values are expressed as a percentage relative to the control condition (means ± SD, n=7-9). Equal loading of Western blots was ensured by β-actin. Means (for the phosphorylated or total protein levels) without a common letter differ ($P<0.05$).

**Figure 2.** Effect of EC and CPE on levels of phosphorylated and total AKT, GSK3 and GS in HepG2 cells. (A) Bands of representative experiments. Percentage data of (B) p-AKT/AKT, (C) p-GSK3/GSK3 and (D) pGS/GS ratios relative to controls. Values are expressed as a percentage relative to the control condition and are means ± SD, n=6-9. Equal loading of Western blots was ensured by β-actin. Means without a common letter differ ($P<0.05$).

**Figure 3.** Effect of EC and CPE on phosphorylated and total AMPK levels in HepG2 cells. (A) Bands of representative experiments. (B) Percentage values of p-AMPK/AMPK ratio relative to the control condition (means ± SD, n=7-8). Equal loading of Western blots was ensured by β-actin. Different letters over bars indicate statistically significant differences ($P<0.05$).
Figure 4. Effect of EC and CPE on GLUT-2 levels in HepG2 cells. (A) Bands of representative experiments. (B) Densitometric quantification of GLUT-2. Values are expressed as a percentage relative to the untreated control condition and are means ± SD, n=7-8. Equal loading of Western blots was ensured by β-actin. Means without a common letter differ (P < 0.05)

Figure 5. Effect of EC and CPE and selective inhibitor LY (LY294002) on levels of PEPCK, AKT and AMPK, and glucose production. HepG2 cells were incubated in the presence or absence of 25 µM LY for 1 h and later with 10 µM EC or 1 µg/mL CPE for 24 h. (A) Bands of representative experiments. Percentage data of (B) PEPCK, (C) p-AKT/AKT and (D) p-AMPK/AMPK relative to the control condition (means ± SD, n=6-9). Equal loading of Western blots was ensured by β-actin. (E) Glucose production was expressed as percent of control are means ± SD of 10-15 different samples per condition. Different letters over bars indicate statistically significant differences (P<0.05).

Figure 6. Effect of EC and CPE and selective inhibitor Compound C (Comp. C) on levels of PEPCK, AKT and AMPK, and glucose production. HepG2 cells were incubated in the presence or absence of 30 µM Comp. C for 1 h and later with 10 µM EC or 1 µg/mL CPE for 24 h. (A) Bands of representative experiments. Densitometric quantification of (B) PEPCK levels, (C) p-AKT/AKT and (D) p-AMPK/AMPK. Values are expressed as a percentage relative to the control condition and are means ± SD, n=6-10. Equal loading of Western blots was ensured by β-actin. (E) Glucose production was expressed as percent of control are means ± SD of 10-16 different samples per condition. Means without a common letter differ (P<0.05).
**Figure 7.** Schematic overview showing the EC- and CPE-induced insulin and glucose signalling pathways analysed in HepG2 cells. Sharp arrows indicate positive inputs (activation), whereas a line shows negative inputs (inhibition).
Table 1. Effects of EC and CPE on (A) cell viability and (B) cell proliferation. Cell viability was determined as relative percent of Crystal Violet stained control cells. Cell proliferation was calculated as percentage of the relative increase over the control values of BrdU incorporated into genomic DNA. Data represent means ± SD of 8-10 samples. Means for each antioxidant without a common letter differ, \( P < 0.05 \).

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<th>Cell viability (% of viable cells)</th>
<th>Cell proliferation (% of controls)</th>
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<td><strong>C</strong></td>
<td>100.5 ± 10.1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>100.0 ± 7.8&lt;sup&gt;a&lt;/sup&gt;</td>
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<td><strong>EC (µM)</strong></td>
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<tr>
<td>1</td>
<td>92.2 ± 6.3&lt;sup&gt;a&lt;/sup&gt;</td>
<td>100.8 ± 9.2&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
<td>5</td>
<td>103.7 ± 5.3&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
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<td>104.7 ± 10.9&lt;sup&gt;a&lt;/sup&gt;</td>
<td>104.1 ± 8.7&lt;sup&gt;a&lt;/sup&gt;</td>
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Figure 3

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<tr>
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<th>P-AMPK/AMPK Optical density (%)</th>
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EC (μM) | CPE (μg/mL) |
---------|-------------|
1        | 1           |
5        | 5           |
10       | 10          |
Figure 4

A

GLUT-2

β-actin

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B

GLUT2 Optical density (%)

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Figure 5

(A) Western blot analysis of PEPCK, p-AKT, AKT, p-AMPK, and AMPK expression in LY, EC, and CPE groups.

(B) Optical density (%) of PEPCK expression in LY, EC, and CPE groups.

(C) Optical density (%) of p-AKT/AKT expression in LY, EC, and CPE groups.

(D) Optical density (%) of p-AMPK/AMPK expression in LY, EC, and CPE groups.

(E) Glucose production (% of control) in LY, EC, and CPE groups.
Figure 7

EC/CPE → IR → IRS-1/2 → AKT → GSK3 → GS

AMPK → PEPCK

Glucose production