UPR-mediated TRIB3 expression correlates with reduced AKT phosphorylation and inability of interleukin 6 to overcome palmitate-induced apoptosis in RINm5F cells

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Abstract

Unfolded protein response (UPR)–mediated pancreatic β-cell death has been described as a common mechanism by which palmitate (PA) and pro-inflammatory cytokines contribute to the development of diabetes. There are evidences that interleukin 6 (IL6) has a protective action against β-cell death induced by pro-inflammatory cytokines; the effects of IL6 on PA-induced apoptosis have not been investigated yet. In the present study, we have demonstrated that PA selectively disrupts IL6-induced RAC-alpha serine/threonine-protein kinase (AKT) activation without interfering with signal transducer and activator of transcription 3 phosphorylation in RINm5F cells. The inability of IL6 to activate AKT in the presence of PA correlated with an inefficient protection against PA-induced apoptosis. In contrast to PA, IL6 efficiently reduced apoptosis induced by pro-inflammatory cytokines. In addition, we have demonstrated that IL6 is unable to overcome PA-stimulated UPR, as assessed by activating transcription factor 4 (ATF4) and C/EBP homologous protein (CHOP) expression, X-box binding protein–1 gene mRNA splicing, and pancreatic eukaryotic initiation factor–2α kinase phosphorylation, whereas no significant induction of UPR by pro-inflammatory cytokines was detected. This unconditional stimulation of UPR and apoptosis by PA was accompanied by the stimulation of CHOP and tribble3 (TRIB3) expression, irrespective of the presence of IL6. These findings suggest that IL6 is unable to protect pancreatic β-cells from PA-induced apoptosis because it does not repress UPR activation. In this way, CHOP and ATF4 might mediate PA-induced TRIB3 expression and, by extension, the suppression of IL6 activation of pro-survival kinase AKT.


Introduction

Obesity and insulin resistance are tightly associated with the onset of type 2 diabetes mellitus (T2DM), possibly due to increased circulating levels of glucose and free fatty acids (FFAs), which are known to induce β-cell apoptosis, a phenomenon known as glucolipotoxicity (Donath et al. 2005). However, the precise relevance and the mechanisms by which FFAs trigger β-cell death are still unclear. In contrast to T2DM, type 1 diabetes mellitus (T1DM) is well defined by pancreatic β-cell injury due to the action of pro-inflammatory cytokines, such as interleukin-1β (IL-1β), interferon-γ (IFN-γ), and tumour necrosis factor–α (TNF–α), which are released by infiltrated immune cells (Cnop et al. 2005).

Several recent studies have described that both pro-inflammatory cytokines and FFAs induce an imbalance between endoplasmic reticulum (ER) folding demand and capacity in the β-cells, the so-called ER stress. The cellular consequence of the ER stress is the unfolded protein response (UPR) activation. The initial goal of UPR is to accomplish ER folding rate to the increased demand. However, if the stress persists, the termination phase of UPR targets apoptosis (Ron & Walter 2007). UPR–activated apoptosis occurs by a yet-unrevealed mechanism, but the increase in c-Jun N-terminal kinase activation and C/EBP homologous protein (CHOP) expression is likely to be the essential step (Kharroubi et al. 2004, Akerfeldt et al. 2008).

Great attention is being paid to IL6 produced by adipocytes due to its possible involvement in the onset of T2DM (Herder et al. 2006). IL6 production is directly proportional to adipose tissue mass, which generates more than 30% of its serum levels (Mohamed-Ali et al. 1997).
Although leading to insulin resistance in peripheral tissues (Glund & Krook 2008), IL6 has been shown to play a complex role in pancreatic β-cell death in diabetes. IL6 was described to counteract pancreatic β-cell apoptosis induced by pro-inflammatory cytokines (Park et al. 2003, Choi et al. 2004), and to increase cell death in primary islets treated with FFAs (Ellingsgaard et al. 2008). Paradoxically, it was observed that the incidence of T1DM in non-obese diabetic (NOD) mice decreased when the mice were crossed with transgenic mice overexpressing IL6 (DiCosmo et al. 1994).

In spite of the evident increased levels of IL6 in obese and insulin-resistant patients and the diabetogenic properties of this cytokine, no data are available regarding the ability of IL6 to modulate FFA-induced ER stress in pancreatic β-cells. In order to assess this issue, we investigated whether IL6 modulates palmitate (PA)-induced β-cell apoptosis by focusing on IL6 intracellular signaling pathways and the main branches of UPR.

Materials and Methods

RINm5F cell culture and incubation

The rat insulinoma cell line RINm5F was cultured in RPMI-1640 medium (11.1 mM glucose and 10% FBS) as described previously (Anhê et al. 2007). For the time-course and dose–response experiments, the cells were incubated for 30 min at 37 °C with Krebs-bicarbonate buffer (KBB) containing 0.3 mM glucose prior to the incubation with rat IL6 (Sigma). For the time-course and dose–response experiments, incubation with IL6 was carried out using KBB at different intervals and at several concentrations (indicated in the figures) respectively. PA (Sigma) treatment at prolonged intervals was carried out in a serum-free medium containing 1% of fatty acid-free BSA. Cells were pre-treated with IL6 (100 ng/ml) for 24 h, and then the medium was replaced with a fresh medium containing IL6 alone or in combination with PA, followed by incubation for an additional 12 h. PA was dissolved in 95% ethanol to a concentration of 100 mM, and heated to 60 °C for 30 min. Alcoholic PA solution was added to the culture medium to yield a final concentration of 0.25% (v/v) was added to all the cultures in which PA was absent.

Treatment with pro-inflammatory cytokine mixture (CM) was carried out in RPMI–1640 medium containing 11·1 mM glucose and 10% FBS. Incubation that was carried out using CM lasted for 24 h, and was preceded by a 24 h incubation with IL6. Thus, in these experiments, the total interval during which the cells were exposed to IL6 was 48 h, and during the last 24 h of this time period, the cells were grown in the presence or absence of CM. CM was prepared using a combination of IL1β (60 U/ml, 4·4 ng/ml), IFN-γ (14 U/ml, 10·3 ng/ml), and TNF-α (185 U/ml, 4·4 ng/ml; Souza et al. 2008).

Transfection of RINm5F cells with small interfering RNA (siRNA) targeted at CHOP (sc-156118, Santa Cruz Biotechnology, Santa Cruz, CA, USA) or scrambled control siRNA (sc-37007, Santa Cruz Biotechnology) was performed as described previously (Anhê et al. 2010). Briefly, the cells were washed twice with a serum-free medium (Opti-MEM, Invitrogen), and were then incubated with 1 ml of the same medium containing the siRNA (final concentration of 80 nM) previously mixed with 2 μl Lipofectamine 2000 (Invitrogen), or Lipofectamine only (CTL). After 7 h, 1 ml of RPMI medium containing 20% FBS and 22·2 mM glucose was added to the culture containing siRNAs and Lipofectamine, and the cells were incubated for an additional 12 h (final concentrations of FBS and glucose were 10% and 11·1 mM respectively). Thereafter, the cells were allowed to grow for 60 h in RPMI medium (11·1 mM glucose and 10% FBS) that was replaced every 24 h. After this period, the cells were treated with PA for 12 h as described above.

Pancreatic islet isolation and culture

Islets were isolated by collagenase digestion as described previously (Bordin et al. 1995). All the experiments involving animals were performed in accordance with the guidelines of the Brazilian College for Animal Experimentation (COBEA). After isolation, groups of ca. 400 islets were extensively and carefully washed with KBB containing antibiotics, and were cultured for 24 h. In order to induce apoptosis, pancreatic islets were cultured with PA in RPMI–1640 medium containing 11·1 mM glucose, 10% FBS, and 2% of fatty acid-free BSA as described previously (Shimabukuro et al. 1998). Total (bound + unbound) PA concentration used was 1 mM, and IL6 concentration was 100 ng/ml. Treatments with PA and/or IL6 were performed for 24 h. After culture, groups of ~250 islets grown under each condition were processed for protein extraction and western blotting or for the detection of DNA fragmentation. For DNA fragmentation, groups of 50 islets were dissociated by re-suspension in Ca2+-free KBB containing EDTA.

Caspase-3 activity

Cells were seeded in six-well plates (5 × 105/well), and treated as described in the figure legends. Afterwards, the cells were centrifuged at 1000 g for 15 min at 4 °C, and 200 μl of a lysis buffer containing 10% sucrose, 0·1% CHAPS, 100 mM HEPES, pH 7·4, 10 mg/ml leupeptin, 10 mg/ml aprotinin, 2 mM phenylmethylsulphonyl fluoride (PMSF), and 10 mM dithiothreitol (DTT) were added to the pellet. The cells were incubated for 30 min at 4 °C and centrifuged at 12 000 g for 30 min at 4 °C. The supernatant was collected, and the protein content was determined by the Bradford method. Caspase-3 protease activity was determined by incubating the lysate (20 μg of total protein) with 50 mM fluorogenic substrate Ac-DEVD-AMC (Sigma) in a buffer containing 10% sucrose, 100 mM HEPES, pH 7·4, 10 mg/ml leupeptin, 10 mg/ml


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aprotinin, 200 mM PMSF, and 10 mM DTT. Caspase-3 activity was assessed by measuring the fluorescence of Asp–7–amino–4-trifluoromethyl coumarin released at 30 min during the 5-min intervals using a spectrofluorometer (Molecular Devices Spectra MAX Gemini XS, Sunnyvale, CA, USA).

**Determination of membrane integrity and DNA fragmentation**

Cells were seeded in six-well plates (5 × 10^5/well), and were treated as described in the figure legends. Afterwards, the cells were harvested with trypsin, washed, and suspended in PBS. Cell suspensions were divided into two aliquots, and were used to determine DNA fragmentation and membrane integrity. Both the methodologies were based on propidium iodide (PI) staining and fluorescence, which was measured using a flow cytometer. First, 50 μl of isotonic solution containing PI (50 mg/ml in PBS) were used to homogenize one pelleted aliquot from each sample, and were immediately analyzed using a FACSCalibur flow cytometer (Becton Dickinson, San Juan, PR, USA). PI is highly hydrosoluble and usually does not significantly diffuse into the cell. However, when membrane integrity is disrupted, it does occur, making the cell emit fluorescence when excited by the laser. Fluorescence was measured using the FL2 channel (orange-red fluorescence; 585/42 nm). Ten thousand events were analyzed per sample using the CellQuest software (Becton Dickinson). DNA fragmentation was measured by staining the second aliquot of the cell suspension obtained from each sample with a PI solution according to the method described previously (Verlengia et al. 2003). In this case, the cells were incubated for 1 h at room temperature (RT) with a PI solution containing 0.1% Triton X–100 (50 mg/ml PI, 0.1% sodium citrate, and 0.1% Triton X–100). The presence of the detergent in the solution makes the cell membrane permeable to PI, allowing it to be incorporated into cellular DNA. Fluorescence was measured and analyzed using the CellQuest software, and low fluorescence was emitted by fragmented DNA, while intact haploid and diploid DNAs emitted characteristic two-peaked high-intensity fluorescence.

**RINm5F cell proliferation**

Cell proliferation was assessed by measuring bromodeoxyuridine (BrdU) incorporation into DNA using a commercial kit (Millipore, Billerica, MA, USA; Cat. no. 2750). Briefly, 4 × 10^4 cells were seeded in a 96–well plate, and were allowed to attach to the wells containing the growth medium overnight. On the next day, the medium was replaced with low serum medium (0.1% FBS), in which each treatment was overnight. On the next day, the medium was replaced with to attach to the wells containing the growth medium emitted characteristic two-peaked high-intensity fluorescence.

**RNA extraction and RT-PCR**

Cells were harvested in TRIzol, and were processed for total RNA extraction and cDNA synthesis, which were performed using conventional (GoTaq DNA Polymerase, Promega) or real-time (SYBR Green master mix, Applied Biosystems, Foster City, CA, USA) PCR. The sequences of the primers and amplification conditions were as follows: B-cell CLL/lymphoma (Bcl2; NM_016993) sense 5′-GCCAGAGATGTCAGTCCAGTCAG-3′ and antisense 5′-CCCACCGA-CTCAAAGAAGG-3′, 128 bp, 55°C; Bcl2 associated X protein (Bax; NM_017059) sense 5′-GGGATGAAGCTC-GCAATAATG-3′ and antisense 5′-GCAAGATGTA-GAAAGGGCAACC-3′, 151 bp, 53°C; β2-microglobulin (β2M; NM_012512) sense 5′-CTCAGTTCCACCACCCTCAG-3′ and antisense 5′-GCAAGCATATACATCGG-3′; 157 bp, 56°C; succinate dehydrogenase subunit A (SdhA; NM_130428) sense 5′-CGGAAGCCCGGAAGGAGT-3′ and antisense 5′-CATAGGAAAGCCCTGAGC-3′, 249 bp, 56°C. Relative expression levels were analyzed using the 2(−ΔΔC(T)) method (Livak & Schmittgen 2001) with β2m and Sdha as the internal controls. Conventional RT-PCR was performed for the analysis of X-box binding protein gene (Xbp1) splicing as described previously (Kharroubi et al. 2004). Briefly, PCR products were incubated with the Pot1 restriction enzyme for 5 h at 37°C, followed by separation on a 2% EtBr agarose gel. Primer sequences and amplification conditions were as follows: Xbp1 (NM_001004210) sense 5′-AACAGAGTAGCAGCGCAGACTGC-3′ and antisense 5′-GGATCTCTAAAATCAGAGGCTTGGTG-3′, 600 bp, 52°C.

**Western blotting**

Cells were harvested in a lysis buffer and homogenized by sonication. After centrifugation, equal amounts of protein were resolved by SDS–PAGE, and transferred onto nitrocellulose membranes. Detection using specific antibodies, HRP-conjugated secondary antibodies, and luminol solution was performed as described previously (Anhê et al. 2006).
Concordantly, PA treatment of RINm5F cells increased in the phosphorylation of both STAT3 and AKT (Fig. 1B). Also assessed, IL6 at 100 ng/ml produced a significant increase of the IL6 receptor, namely STAT3 and protein kinase B/AKT. The phosphorylation of both STAT3-tyrosine and AKT-serine peaked at 15 min after incubation with IL6 (Fig. 1A). The concentration of IL6 used in the further experiments was also assessed. IL6 at 100 ng/ml produced a significant increase in the phosphorylation of both STAT3 and AKT (Fig. 1B). Figure 1C shows that CM-treated cells displayed higher DNA fragmentation (2.3-fold of the CTL values) which was paralleled by CM-induced caspase-3 activity (10.8-fold of the CTL values). Figure 1D shows that CM increased caspase-3 activity (10.8-fold of the CTL values). Afterwards, the cells were processed for protein extraction and detection of STAT3, pSTAT3(Tyr), AKT, and pAKT(Ser) (n = 3). RINm5F cells were treated with IL6 (100 ng/ml) in combination with or without a mixture of IL1β (4-4 ng/ml), IFN-γ (10-3 ng/ml), and TNF-α (4-4 ng/ml); cytokine mixture, CM). Untreated cells are designated as CTL cells. After 24 h, cellular DNA fragmentation (C), caspase-3 activity (D), cell membrane integrity (E), and cell proliferation rate (F) were analyzed (see Materials and Methods). Results are shown as mean ± S.E.M. Comparisons were performed using one-way ANOVA with the Tukey-Kramer post-test (InStat – GraphPad Software, Inc., San Diego, CA, USA). The level of significance was set at P<0.05.

Results

IL6 modulation of STAT3 and AKT phosphorylation and pro-inflammatory cytokine-induced apoptosis and suppression of proliferation

We have demonstrated the activation of proteins downstream of the IL6 receptor, namely STAT3 and protein kinase B/AKT. The phosphorylation of both STAT3-tyrosine and AKT-serine peaked at 15 min after incubation with IL6 (Fig. 1A). Figure 1C shows that CM-treated cells displayed higher DNA fragmentation (2.3-fold of the CTL values) which was abolished by pre-treatment with IL6. These changes paralleled the modulation of caspase-3 activity. Figure 1D shows that CM increased caspase-3 activity (10.8-fold of the CTL values) and IL6 diminished CM-induced caspase-3 activation (0.36-fold compared with the CM-treated cells). Cell viability was not significantly changed by CM treatment (Fig. 1E). In contrast, Fig. 1F shows that both IL6 and CM decreased cell proliferation (to 0.85- and to 0.56-fold of the CTL values respectively). Additionally, the combination of CM and IL6 prompted a more pronounced reduction in cell proliferation (0.03-fold of the CTL values).

Effects of IL6 on PA-induced cell death and proliferation, and STAT3 and AKT phosphorylation

The percentage of RINm5F cells with fragmented DNA increased approximately fourfold on PA treatment (Fig. 2A). Concordantly, PA treatment of RINm5F cells increased caspase-3 activity ∼ 30-fold when compared with the CTL cells (Fig. 2B). IL6 treatment did not change basal or PA-induced DNA fragmentation and caspase-3 activation. PA treatment reduced cell viability to ∼ 0.8-fold of the CTL values, and this effect was not modulated by IL6 treatment (Fig. 2C). Figure 2D shows that both IL6 and PA decreased cell proliferation (to 0.85- and to 0.37-fold of the CTL values respectively). The combination of IL6 and PA did not promote a further reduction in cell proliferation.

On the other hand, IL6 signaling was differentially regulated by PA. The phosphorylation of both STAT3 and AKT was increased by IL6 (2.1-fold of the CTL values for both STAT3 and AKT), but only IL6-induced AKT phosphorylation was suppressed by PA (Fig. 3).

Regulation of pro- and anti-apoptotic proteins by IL6 and PA

IL6 by itself increased the expression of the anti-apoptotic Bcl2 mRNA (2.7-fold of the CTL values). Bcl2 mRNA expression was suppressed by PA (Fig. 4).
expression was downregulated by PA alone or in combination with IL6 (by 0.5- and 0.4-fold of the CTL values respectively; Fig. 4A). The absolute levels of BCL2 protein were not altered by either IL6 or PA (Fig. 4B). The levels of the pro-apoptotic BCL2-associated X protein (Bax) mRNA were downregulated by IL6 (0.3-fold of the CTL values), and upregulated by PA alone or in combination with IL6 (by 2.2- and 1.8-fold of the CTL values respectively; Fig. 4C). IL6 decreased BAX protein content (0.6-fold of the CTL values), but the presence of PA abolished this effect (Fig. 4D). IL6 increased Bcl2/Bax mRNA ratio (10-fold of the CTL values), whereas PA decreased it (0.3-fold of the CTL values). In addition, Bcl2/Bax mRNA ratio remained reduced when IL6 was combined with PA (Fig. 4D). BCL2/BAX protein ratio was increased by IL6 alone (1.8-fold of the CTL values), but this effect was abolished when PA was added to the medium (Fig. 4F).

**Analysis of UPR and TRB3 expression under PA and IL6 treatments**

Representative blots of the effects of IL6 and PA treatments on BiP, ATF4, CHOP, and TRIB3 protein contents are shown in Fig. 5A. Both the appearance of phosphorylated PERK (Fig. 5B) and the splicing of Xbp1 (Fig. 5C) were stimulated by PA, in such a way that the addition of IL6 did not interfere with this upregulation. Neither IL6 nor PA modulated BiP content (Fig. 5D). On the other hand, ATF4 and CHOP contents were increased by PA treatment (1.9- and 2.2-fold of the CTL values respectively; Fig. 5E and F respectively). Addition of IL6 to the medium did not modulate PA-induced increase in ATF4 and CHOP. PA, either alone or in combination with IL6, did not change eIF2α phosphorylation (data not shown). We have also shown that the expression of TRIB3 presented a pattern that was the
shown as mean BCL2 and BAX protein content were also calculated (F). Results are (B) and BAX (D). The ratio between processed for protein extraction and western blot detection of BCL2 Another set of cells was subjected to the same treatment, and processed for RNA extraction and cDNA synthesis. CTL cells were treated with a vehicle. Samples were used for real-time PCR analysis of cDNA synthesis. CTL cells were treated with a vehicle. Samples were used for real-time PCR analysis of Bcl2 / Bax (C). The intensity of the bands of BiP (D), ATF4 (E), CHOP and TRIB3 protein contents were not changed by CM and/or IL6 treatment. same as that of ATF4 and CHOP. TRIB3 protein levels were increased by PA treatment (1.8-fold of the CTL values), and was the same as that found for TRIB3 protein expression was detected for Trib3 mRNA expression analyzed by real-time PCR (data not shown).

**Modulation of DNA fragmentation and ER stress by IL6 in rat pancreatic islets cultured with PA**

Our data show that both IL6 and PA increase DNA fragmentation in pancreatic islets cultured for 24 h (to ~1-6-fold of the CTL values). The combination of PA and IL6 resulted in a further increase in DNA fragmentation (to 2-4-fold of the CTL values; Fig. 6A). Figure 6B shows the representative blots of CHOP and TRIB3 obtained in cultured pancreatic islets. PA increased both CHOP and TRIB3 protein contents in islets to 2-4- and 3-0-fold of the CTL values respectively. These changes were not modulated by IL6 (Fig. 6C and D respectively).

**Analysis of UPR and TRIB3 expression under CM and IL6 treatments**

Representative blots of the effects of IL6 and CM treatments on BiP, ATF4, CHOP and TRIB3 protein contents are shown in Fig. 7A. Neither IL6 nor CM modulated BiP content (Fig. 7B). CM induced a 1-6-fold increase in ATF4 content, which was suppressed by the presence of IL6 (Fig. 7C). In addition, CHOP (Fig. 7D) and TRIB3 (Fig. 7E) contents were not changed by CM and/or IL6 treatment.

**CHOP knockdown with siRNA suppresses PA-induced apoptosis in RINm5F cells**

As shown previously herein, PA induced an increase in DNA fragmentation in RINm5F cells (Fig. 8B). Specific knockdown of CHOP with siRNA (Fig. 8A) had no effect on basal DNA fragmentation, but it abolished the ability of PA to increase it. DNA fragmentation that was induced by PA in
RINm5F cells transfected with siRNA targeted at CHOP was 0.7-fold of that induced by PA in untransfected cells, reaching values similar to the CTL values (1.1-fold).

**Discussion**

IL6 has been suggested to protect pancreatic β-cells from apoptosis induced by TNF-α, IFN-γ, and IL-1β, which are the main pro-inflammatory cytokines involved in the assault of the endocrine pancreas during the onset of T1DM (Park et al. 2003, Choi et al. 2004). Moreover, mating of mice that overexpress IL6 with NOD mice decreased the incidence of diabetes (DiCosmo et al. 1994). On the other hand, IL6 was reported to decrease insulin secretion and β-cell proliferation (Sandler et al. 1990). These observations show that the role of IL6 in β-cell survival and function is complex, and justify additional studies in order to clarify the intracellular mechanisms and cellular responses elicited by this cytokine in pancreatic β-cells. To further address this issue, we sought to investigate if IL6 protects β-cell against lipotoxicity induced by FFAs.

Our study has demonstrated that IL6 activates its classical target STAT3 in RINm5F cells. AKT pathway, a non-canonical signaling pathway of IL6 (Hsu et al. 2004), was also activated in RINm5F cells. IL6 by itself enhanced Bcl2 expression and decreased Bax expression, probably as a result of STAT3 activation (Kovalovich et al. 2001). Inhibition of the mitochondrial pathway of apoptosis by adipokines has been documented already and is in agreement with our results. Leptin, which shares intracellular signaling pathways with IL6, increases Bcl2 and decreases BAX, altering the BCL2:BAX ratio and reducing apoptosis induced by serum depletion in BRIN-BD11 β-cell line (Brown & Dunmore 2007). Furthermore, we observed that IL6 decreased apoptosis induced by pro-inflammatory cytokines. Altogether, our results confirm that IL6 protects RINm5F cells against pro-inflammatory cytokine-induced apoptosis, as has been published already by others (Park et al. 2003, Choi et al. 2004). In contrast, we have shown that IL6 potentiates the reduction of pancreatic β-cell proliferation induced by pro-inflammatory cytokines. We currently have no data to explain this finding. A hypothesis is that IL6 promotes the growth arrest of RINm5F cells, as it does in other cell types (Kamimura et al. 2003). Of note, our experiments were carried out for a short period during which IL6 treatment was given, which was described to decrease islet DNA content (Sandler et al. 1990).

![Figure 6](image)

**Figure 6** Effects of palmitate and IL6 on apoptosis and UPR-related proteins in pancreatic rat islets. Isolated rat islets were treated with IL6 (100 ng/ml) and palmitate (PA; 1 mM) for 24 h, and processed for the analysis of DNA fragmentation by propidium iodide fluorescence (A) and western blotting. In (B), the representative film image of CHOP, TRIB3, and β-actin (loading control) is shown. In (C) and (D), the mean±s.e.m. of CHOP and TRIB3 protein contents that were obtained from two to three independent experiments and analyzed by densitometry is shown. *P<0.05 versus CTL; and **P<0.01 and ***P<0.001 versus CTL and IL6 (n=6–18 for DNA fragmentation analysis).

![Figure 7](image)

**Figure 7** Modulation of UPR-related proteins by cytokines mixture and IL6. RINm5F cells were treated with IL6 (100 ng/ml) in combination with or without a mixture of IL1β (4–4 ng/ml), IFN-γ (10–3 ng/ml), and TNFα (4–4 ng/ml); cytokine mixture, CM. CTL cells were treated with a vehicle. After 24 h, the cells were processed for protein extraction and western blot detection of BiP, ATF4, CHOP, TRIB3, and the internal control β-actin (A). The intensity of the bands of BiP (B), ATF4 (C), CHOP (D), and TRIB3 (E) was analyzed by densitometry using different sets of experiments. Results are shown as mean±s.e.m. *P<0.05 versus CTL (n=4–6).
Figure 8 Effect of palmitate on DNA fragmentation after knockdown of CHOP with siRNA. RINm5F cells were transfected with small interfering RNA targeted at CHOP (CHOP siRNA) or with a scrambled control siRNA (scrambled siRNA). Lipofectamine 2000 was mixed with siRNAs and used in the CTL experiment. Palmitate (PA; 250 μM) was added during the last 12 h of the 72-h period after the transfection; after that, the cells were processed for total protein extraction and SDS-PAGE. Samples from CTL, scrambled siRNA, and CHOP siRNA cells were subjected to immunodetection of CHOP and β-actin, the latter being used to check the specificity of CHOP siRNA (A). DNA fragmentation was analyzed by propidium iodide fluorescence assessed using a flow cytometer (B). ***P<0.001 versus CTL and *p<0.001 versus PA (n=7–15).

Pancreatic β-cell death occurs in both T1DM and T2DM, but in opposition to that in T1DM, β-cell apoptosis in T2DM is likely to result mainly from lipotoxicity and glucotoxicity. PA, are of the most abundant FFAs found in serum (Hamilton & Kamp 1999), is known as a potent inducer of β-cell apoptosis in primary β-cells and RINm5F cells (Beeharry et al. 2004, Azevedo-Martins et al. 2006). Our results show that IL6 did not protect RINm5F cells from PA-induced apoptosis. This finding is supported by clinical observations showing that obese and insulin-resistant patients often develop T2DM regardless of increased IL6 circulating levels (Herder et al. 2006).

In contrast to that observed in β-cell lineage, our results show that IL6 potentiated PA-induced apoptosis in cells obtained from cultured islets. Ellingsgaard et al. (2008) found similar results using cultured mouse islets. We do not have any explanation for this, but we can speculate that the release of cytotoxic mediators by the variety of resident cell types in islets, that is, macrophages, endothelial cells, and endocrine cells, could play a detrimental role in islet survival. Of note, collagenase-isolated islets have higher apoptotic rate than β-cell lineages, probably due, at least in part, to the release of pro-inflammatory cytokines induced by endotoxin contamination of the conventional type V collagenase (Berney et al. 2001). Indeed, several reports have shown that IL6 alone is not cytotoxic to β-cells (reviewed by Kristiansen & Mandrup-Poulsen (2005)). However, phenotypic differences between primary β-cells and clonal β-cell lines cannot be ruled out.

The mechanism underlying the inability of IL6 to protect RINm5F cells against PA-induced apoptosis probably relies on the disruption of IL6 signaling pathways. We have found that phosphorylation of both STAT3 and AKT was activated by IL6, but only that of AKT was impaired by PA. In pancreatic β-cells, downregulation of STAT3 activity is mainly involved in the modulation of insulin secretion (Gorogawa et al. 2004, Anhê et al. 2007). On the other hand, it is well documented that the phosphatidylinositol 3-kinase (PI3K)/AKT signaling is a critical pathway regulating β-cell survival (Elghazi et al. 2007), which is impaired by FFAs (Wrede et al. 2002, Kharroubi et al. 2004). Downstream to AKT activation, a myriad of pro-apoptotic proteins can be phosphorylated and inhibited, such as BCL2 family members, GSK3, and FoxO1, resulting in a relief from apoptosis (Datta et al. 1999, Wrede et al. 2002, Martinez et al. 2008). Importantly, overexpression of a constitutively active AKT prevented FFA-induced apoptosis (Wrede et al. 2002).

ER stress-activated UPR has been shown to occur in pancreatic islets obtained from diabetic obese animals and human patients with T2DM (Laybutt et al. 2007). Excess of saturated FFAs can directly trigger ER stress and UPR-mediated apoptosis in the pancreatic β-cell-derived lineages MIN6 (Laybutt et al. 2007) and INS1 (Karasov et al. 2006) and primary rat pancreatic β-cells (Kharroubi et al. 2004). Here, we have shown that UPR was activated in RINm5F cells by PA after 12-h incubation (250 μM), i.e. in a shorter period and at a lower dose than those required for MIN6 and INS1. Therefore, RINm5F cells are likely to be more sensitive to PA-induced ER stress, in spite of there being no change in BiP expression and eIF2α phosphorylation. These later effects were not unexpected, since it has been previously demonstrated that, in INS1 cells, BiP levels are not affected by PA (Karasov et al. 2006, Diakogiannaki et al. 2008). Moreover, eIF2α phosphorylation is likely to be transient, returning to basal levels after 6–16 h of treatment. Our results confirm that UPR is far from being an all-or-nothing response, and that it is differentially activated according to the cell type and the stressor.

UPR consists of three branches of signaling pathways that initially aim to adjust the increased ER demand. Activation of UPR is triggered by the dissociation of the chaperone BiP from the transducer proteins PERK, ATF6, and IRE1. BiP dissociation from PERK results in its activation that leads to eIF2α phosphorylation and ATF4 expression. This initial pathway inhibits general translation rate, and therefore, decreases the flux of new synthesized proteins to the ER (Harding & Ron 2002). Thus, disruption of the
PERK/eIF2α/ATF4 pathway results in the inability of pancreatic β-cells to deal with increased folding demand, and therefore, accelerates ER stress-mediated apoptosis (Yamaguchi et al. 2008). In spite of the protective feature of this UPR branch, PA has been shown to stimulate the PERK/eIF2α/ATF4 pathway prior to the induction of pancreatic β-cell death (Kharroubi et al. 2004). Herein, we have shown that the lack of IL6 protection against PA-induced β-cell apoptosis correlates with the inability of this IL to prevent PERK phosphorylation and ATF4 expression induced by PA, reinforcing the idea that UPR might mediate lipotoxicity in RINm5F cells.

Similarly, IL6 did not modulate PA-induced increase in Xbp1 mRNA splicing. Splicing of Xbp1 mRNA is catalyzed by IRE1α under conditions of ER disturbance. Spliced mRNA of Xbp1 encodes for an active transcription factor that facilitates the expression of genes involved in the ER protein degradation (Harding & Ron 2002). The spliced mRNA of Xbp1 has been shown to be increased in the pancreatic islets obtained from diabetic mice, and in the pancreatic islets treated with PA or inflammatory cytokines (Karaskov et al. 2006, Akerfeldt et al. 2008).

We and others have demonstrated that PA increases CHOP expression in pancreatic β-cells (Wang et al. 2005, Karaskov et al. 2006, Laybutt et al. 2007, Martinez et al. 2008). CHOP is regulated by ATF4, and plays an essential role in the UPR-mediated apoptosis in pancreatic β-cells (Pirot et al. 2007) in such a way that CHOP deletion improves pancreatic β-cell function in diabetic rodents (Song et al. 2008). As observed for ATF4, IL6 was not able to modulate the increase in CHOP induced by PA. The participation of CHOP in the induction of apoptosis is not completely understood, but one hypothesis is that this transcription factor may suppress Bcl2 expression (McCullough et al. 2001). In accordance with ATF4 and CHOP results, the decrease in Bcl2 expression induced by PA was not changed by IL6. Reinforcing the role of CHOP in FFA-induced apoptosis, we have shown that siRNA-mediated knockdown of CHOP abolishes PA-induced apoptosis in RINm5F cells. This result matches the data that have been published already by performing similar experiments using INS-1 cells (Akerfeldt et al. 2008).

Another important mechanism by which CHOP links ER stress to apoptosis is the induction of TRIB3 expression after dimerization with ATF4 (Ohoka et al. 2005). Herein, we have shown that PA increases TRIB3 content irrespective of the presence of IL6. As TRIB3 has been described to function as a pseudokinase that inhibits AKT (Du et al. 2003), it is likely that PA-induced TRIB3 expression inhibits AKT-serine phosphorylation stimulated by IL6. In accordance with our results, both PA and high glucose concentrations can trigger UPR-dependent apoptosis in pancreatic β-cells and concurrently increase TRIB3 expression (Martinez et al. 2008, Qian et al. 2008).

Our results further suggest that the modulation of the BCL2:BAX ratio by PA in RINm5F cells is possibly linked to the activation of ER stress. This can be hypothesized because i) CHOP was demonstrated to downregulate BCL2 expression (McCullough et al. 2001), and ii) reduced AKT activation might allow the transcription factor FoxO1 to translocate to the nucleus and to induce BAX expression (Kim et al. 2005). In the latter mechanism, ER stress can disrupt AKT activation by inducing TRIB3 expression. In accordance with these two mechanisms, our results show concordant suppression of IL6-induced Bcl2 expression and increase in CHOP by PA. We have also shown that PA sustains high levels of Bax expression in parallel to increased TRIB3, and decreases AKT activity.

Akerfeldt et al. have demonstrated that although pro-inflammatory ILs are able to induce ER stress and activate UPR, this mechanism is not required for the pro-apoptotic action of these cytokines. On the other hand, these authors suggested that the activation of UPR is a determinant event that culminates in the apoptosis induced by PA (Akerfeldt et al. 2008). Considering these observations, our data suggest that IL6 does not prevent or abolish the disturbance in ER homeostasis elicited by PA, and thus it is not able to protect pancreatic β-cells from apoptosis induced by elevated FFAs. In contrast, as we and others have demonstrated, IL6 can properly protect β-cells against pro-inflammatory-induced apoptosis probably because in this case it has been consistently demonstrated that ER stress does not mediate cell death.

Corroborating this hypothesis, we have shown that the combination of pro-inflammatory cytokines used by us efficiently increases DNA fragmentation and caspase-3 activity without increasing BiP, TRIB3, and CHOP, which are proteins that link ER stress to apoptosis in pancreatic β-cells. In contrast, Akerfeldt et al. (2008) reported that inflammatory cytokines are able to increase CHOP and BiP expression. One plausible explanation for these differences is that the mixture of inflammatory cytokines used by Akerfeldt contained IFN-γ at a concentration that was ten times that of IFN-γ that was used by us. In our experimental conditions, the only UPR-related protein that paralleled the increase in apoptosis induced by CM in RINm5F cells was ATF4, which was suppressed by IL6. The exact role of this modulation is not clear so far, since the PERK/eIF2α/ATF4 pathway seems to be protective rather than pro-apoptotic to pancreatic β-cells (Yamaguchi et al. 2008).

Taken together, our results show that differently from the pro-inflammatory cytokines that trigger apoptosis, IL6 does not prevent PA-induced β-cell death. This effect is probably due to the inability of IL6 to repress UPR activation and TRIB3 expression, which may result in the blockage of AKT activation.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.
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