RFamide peptides 43RFa and 26RFa both promote survival of pancreatic β-cells and human pancreatic islets but exert opposite effects on insulin secretion.
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**Running title:** Effects of 43RFα and 26RFα in pancreatic β-cells

**Word count:** 4892

**Number of Figures:** 6

**Number of Tables:** 0
Abstract

RFamide peptides 43RFa and 26RFa have been shown to promote food intake and to exert different peripheral actions through GPR103 receptor binding. Moreover, 26RFa was found to inhibit pancreatic insulin secretion, whereas the role of 43RFa on β-cell function is unknown, as well as the effects of both peptides on β-cell survival. Herein, we investigated the effects of 43RFa and 26RFa on survival and apoptosis of pancreatic β-cells and human pancreatic islets. In addition, we explored the role of these peptides on insulin secretion and the underlying signaling mechanisms. Our results show that in INS-1E β-cells and human pancreatic islets, both 43RFa and 26RFa prevented cell death and apoptosis induced by serum starvation, cytokine synergism and glucolipotoxicity, through PI3K/Akt- and ERK1/2-mediated signaling. Moreover, 43RFa promoted, whereas 26RFa inhibited glucose- and exendin-4-induced insulin secretion, through Gαs and Gαi/o proteins, respectively. Inhibition of GPR103 expression by small interfering RNA blocked 43RFa insulinotropic effect, but not the insulinostatic action of 26RFa. Finally, 43RFa, but not 26RFa, induced cAMP increase and glucose uptake. In conclusion, because of their survival effects along with the effects on insulin secretion, these findings suggest potential for 43RFa and 26RFa as therapeutic targets in the treatment of diabetes.
Pancreatic beta cell mass plays an essential role in glucose homeostasis. The reduced capacity of the endocrine pancreas to maintain an adequate insulin secretion, due to decreased β-cell mass and function, underlies both type 1 and type 2 diabetes (1). In type 1 diabetes, immune-mediated release of inflammatory cytokines such as tumor necrosis factor-alpha (TNF-α), interferon-gamma (IFN-γ) and interleukin-1β (IL-1β) has been implicated in β-cell apoptosis (2). In type 2 diabetes, β-cell apoptosis results from the combined action of increased plasma glucose/free fatty acid levels (glucolipotoxicity) (3) and cytokines (4). Therefore, identifying molecules capable of increasing pancreatic β-cell survival may be crucial for the treatment and prevention of diabetes.

RFamide-related peptides constitute a family of biologically active peptides terminating in arginine-phenylalanine-amide (Arg-Phe-NH₂) at their C-terminus. They include a 26-aminoacid RFamide peptide (26RFa), that was isolated from the frog brain (5), and a longer form of 26RFa, a 43-aminoacid peptide with a pyroglutamylated RFamide (termed QRFPO43 or 43RFa). Both peptides were identified as the cognate ligands of the orphan G protein-coupled receptor 103 (GPR103) (6,7). In humans and rodents, 43RFa, 26RFa and GPR103 are mainly expressed in brain, particularly in the hypothalamus (8), as well as in peripheral tissues, including the eye, testis, thyroid, adipose tissue and macrophages (6,9-11). 43RFa and 26RFa have been implicated in many physiological functions, including stimulation of food intake (5,12-17), regulation of gonadotropic axis (18,19), aldosterone secretion (7,20), bone formation (21), adipogenesis and inflammation (10,11), blood pressure (12) and prostate cancer differentiation and migration (22).

Recently, 26RFa has been shown to regulate peripheral glucose metabolism through inhibition of glucose-induced insulin secretion in the perfused rat pancreas, without affecting glucagon secretion. This effect involved inhibition of the adenylyl cyclase/cAMP system via a pertussin toxin (PTX)-sensitive Gi protein (23). On the other hand, to our knowledge, the role of 43RFa on insulin secretion is unknown, as well as the effect of both peptides on pancreatic β-cell survival. Therefore, in the present study we sought to determine the role of both 43RFa and 26RFa on survival and apoptosis of pancreatic β-cells and human pancreatic islets, particularly in stress conditions such as
serum starvation, cytokine synergism or glucolipotoxicity. Furthermore, we investigated the effects of the peptides on insulin secretion and the underlying signaling mechanisms.
Research Design and Methods

Reagents

Rat QRFP-43 (43RFa), QRFP-26 (26RFa) and Exendin-4 were from Phoenix Pharmaceuticals (Karlsruhe, Germany). PD-98059, Wortmannin, PTX, NF449, Hoechst-33258, 3-isobutyl-1-methylxanthine (IBMX), MTT, forskolin, glucose, palmitate, Mammalian cell lysis kit were from Sigma-Aldrich (Milano, Italy). Cytokines (TNF-α, INF-γ and IL-1β) were from Life Technologies, Milan Italy. [3H]-2-deoxi-D-glucose was from Perkin Elmer Life Sciences Inc. (Boston, MA). Cell culture and RT-PCR reagents were from Life Technologies (Milan, Italy). P-Akt (Ser473), P-ERK1/2, ERK1/2 and Akt antibodies were from Cell Signaling Technology (Euroclone, Milan, Italy), β-actin antibody was from Santa Cruz (D.B.A. Italia, Milan, Italy). GPR103-antibody was from Abcam (Prodotti Gianni, Italy) and QRFP from Phoenix Pharmaceuticals (Burlingame, CA), for either immunohistochemistry, immunofluorescence and Western blot experiments. Primers for RT-PCR were from from IDT (Tema Ricerca, Bologna, Italy).

Cell culture

INS-1E rat β-cells were kindly provided by Prof. Claes B. Wollheim (Department of Cell Physiology and Metabolism, University of Geneva, Geneva, Switzerland) and cultured as described (24).

Human islet isolation

Human islets were obtained from pancreases of multiorgan donors as described (24). Islet preparations with purity >70%, not suitable for transplantation, were used after approval by the local ethical committee. Islets (10,000) were cultured in CMRL (Invitrogen) with 10% FBS.

RT-PCR and Real Time PCR
Total RNA extraction and reversed transcription to cDNA from 3 µg RNA was performed as described (24). The following primer sequences were used: rat/mouse/human GPR103, Fwd: 5’-TAGGATCAACCATGTCGACGT-3’, Rev: 5’-AAGAGAGCCACCACCTGTCACCATC-3’ [AB109629.1] (21); human QRFP, Fwd 5’-ATGGTAAGCGCTTACCCCCTGATCTAC-3’, Rev 5’-CCTGGAGCTGTGAAGCTCT-3’ (BC101127.2); rat QRFP, Fwd 5’-AGCACACTGGGCTCTCCGTCTAG-3’, Rev 5’-CGCTGGAGCTCTCGAGTCAAGAGTCA-3’ (NM198200.1) (7); 18S rRNA, Fwd: 5’-GTGGAGATTTGTCTGGTT-3’, Rev: 5’-CGCTGAGCCAGTCAGTGTA-3’ (X_01117). 9 µl cDNA were amplified (GeneAmp PCR System, Perkin Elmer, Milan, Italy) in 50 µl under the following conditions: 94°C for 30 sec, 60°C for 30 sec annealing; 72°C for 60 sec, 72°C for 7 min. The final PCR products (318 bp for GPR103, 196 bp for human QRFP, 300 bp for rat QRFP, 199 bp for 18s rRNA) were separated by 1.5% agarose gel electrophoresis and visualized by ethidium bromide staining.

**GPR103 immunofluorescence**

INS-1E β-cells and human pancreatic islets were seeded on glass coverslips in complete medium for 2 days, then washed twice with PBS. INS-1E cells were allowed to grow up to 50-60% confluence. The cells were then fixed with 4% paraformaldehyde and incubated with goat anti-GPR103 antibody overnight at 4°C. After washing, three times with PBS, cells were incubated with anti-goat secondary TRITC-conjugated antibody (Life Technologies, Milan, Italy) for 1 h at room temperature. Nuclei were stained with 4’-6-diamidino-2-phenylindole (DAPI). Red and blue channels were assigned to GPR103 and nuclei, respectively. Images were taken using a Leica DM200 fluorescent microscope and a Leica DFC340 FX camera. Adult rat hippocampal (AHP) cells were used as positive control.

**Immunohistochemistry**
Human pancreatic tissue (n=3 samples) was obtained from surgical samples of non-pancreatic diseases; pituitary samples (n=3 samples) were obtained from autopsies. Tissues were fixed in buffered formalin and embedded in paraffin. Five-µm-thick sections were de-paraffinized and re-hydrated in graded alcohols and phosphate-buffer pH 7.5, and endogenous peroxidase activity was blocked by absolute methanol and 0.3% hydrogen peroxide for 15 min. To assess the presence of GPR103 and QRFP proteins, the following primary antibodies and conditions were employed: rabbit polyclonal antibody to GPR103 (Abcam, Cambridge, UK; diluted 1:150) and rabbit polyclonal antibody to QRFP-43 (Phoenix Pharmaceuticals, Burlingame CA, USA; diluted 1:150). Immunoreactions were revealed by a dextran-chain (biotin-free) detection system (EnVision; Dako), using 3,3-diaminobenzidine (DAB; Dako) as a chromogen.

**Cell proliferation and survival**

Cell proliferation was assessed by BrdU incorporation ELISA (Roche), as previously described (25). Cells were seeded on 96-well plates at 5x10³ cells/well in serum-containing medium until 60–70% confluence and serum-starved for 24 h before treatments. Cell survival was assessed by MTT as previously described (25).

**Hoechst staining**

Morphological changes in the nuclear chromatin of apoptotic cells were detected by Hoechst 33258 staining as previously described (24). 500 stained nuclei were double counted under a fluorescence microscope (DAPI filter).

**Caspase 3 activity**

Caspase 3 activity was assessed by Caspase-3 Colorimetric Kit (Assay Designs, Italy) in INS-1E β-cells and human islet cell lysates, according to the manufacturer’s instruction.
cAMP assay
Starved 8X10^5 INS-1E β-cells and 1X10^3 human islets were seeded in 100-mm dishes. After incubations, in the presence of IBMX (100 µmol/l), cAMP was measured from lysates, using the Direct Cyclic AMP EIA kit (Assay Designs, Italy) according to the manufacturer’s instructions. Forskolin was used as positive control.

Western blotting
40 µg proteins for P-ERK and P-Akt, and 60 µg for GPR103 and QRFP were resolved in 12% SDS-PAGE. Proteins were treated as described (24) and incubated with the specific antibody (dilution 1:1000). Blots were reprobed with the respective total antibodies or with β-actin for normalization. Immunoreactive proteins were visualized with Chemidoc XRS (Bio-Rad, Milan Italy), and densitometric analysis was performed with Quantity One software (Bio-Rad).

Insulin secretion
Insulin secretion in INS-1E β-cells was performed as described (24). Human pancreatic islets (n=3) were incubated for 1 h at 37 C in HEPES-buffered Krebs-Ringer bicarbonate buffer containing 0.5% BSA with 2 mmol/L glucose. The medium was changed, and the cells were incubated again for 1 h in Krebs-Ringer bicarbonate buffer/0.5% BSA containing 1.25, 7.5, 15 or 25 mmol/L glucose, with or without the different peptides or inhibitors. After acid ethanol extraction of the hormone, secreted insulin was quantified by a RIA kit (Linco Research, Labodia, Yens, Switzerland) that recognizes human insulin and cross-reacts with rat insulin.

Small interfering RNA (siRNA)
Rat Qrfpr Silencer Select Pre-designed siRNA (s160302) (GPR103 siRNA), Silencer Negative Control siRNA #1 (cat AM4611) and Lipofectamine RNAiMAX were purchased from Life Technologies (Monza, Italy). INS-1E cells (2 d post seeding) were transfected in RPMI-1640 serum
free medium (Sigma-Aldrich) without antibiotics, with 50 nmol/L Control or GPR103 siRNAs using Lipofectamine RNAiMAX, according to the manufacturer's instructions. After 24 h the cells were used for insulin measurement experiments. Efficiency of transfection was assessed by RT-PCR.

[^H]2-deoxyglucose uptake

[^H]2-deoxyglucose uptake was performed in both INS-1E β-cells and human pancreatic islets. Pancreatic islets were dissociated with StemProAccutase (Gibco, Invitrogen, San Diego, CA, USA) (25). β-cells were incubated in Heps buffer without glucose, with either Ex-4, 43RFa or 26RFa for 20 min, and then with 6 mmol/L 2-deoxyglucose and 1 Ci/mL[^H]-2-deoxyglucose for 5 or 10 min, without Heps buffer removal. Human islets were incubated in serum-free medium with the peptides for 20 min, then the media was aspirated and Heps buffer containing 20 mmol/L 2-deoxyglucose and 1 Ci/mL[^H]-2-deoxyglucose was added for 10 min. The reaction was stopped by washing 3 times with ice cold PBS. Cells were solubilized on ice with lysis buffer (NaOH 50 mmol/L). Equal amounts of lysates were mixed with scintillation fluid, and specific activity was counted in a liquid scintillation counter (Beckman Instruments, Fullerton, CA) in duplicate.[^H]-2-deoxyglucose uptake, expressed as pmol//mg, was normalized by using a BCA protein quantification of each sample, as described (26).

Statistical analysis

Results are presented as means ± SE. Results were analyzed using 2-tailed Student’s t test or two-way ANOVA followed by Tukey’s HSD for post-ANOVA comparisons (GraphPad Prism 5.0 software, San Diego, CA). Significance was established when P<0.05.
Results

GPR103 and QRFP are expressed in INS1-E β-cells and human pancreatic islets

RT-PCR analysis showed mRNA expression of both GPR103 and QRFP, the precursor of RF43a and 26RFa, in pancreatic INS1-E β-cells and human pancreatic islets (Figure 1A and B). At the protein level, Western blot analysis showed presence of both GPR103 and QRFP, the precursor peptide of RF43a and 26RFa, in INS-1E β-cells and human pancreatic islets (Figure 1C and D). Furthermore, immunofluorescence analysis revealed GPR103 protein expression in both β-cells (Figure 1E) and human islets, where GPR103 showed partial colocalization with insulin in β-cells (Figure 1F). Immunohistochemical studies also confirmed expression of both GPR103 and QRFP, in INS1-E β-cells and human pancreatic islets (Figure 1G and H). In human pancreas, GPR103 positivity was found in endocrine pancreatic cells, being negative in acinar cells. Similarly, QRFP was positive in pancreatic islets and negative in acinar cells (Figure 1H). Pituitary was used as positive control, showing strong staining for GPR103 in the posterior lobe (neurohypophysis) but almost negative staining in the anterior lobe (adenohypophysis) (Figure 1H). QRFP was negative in the neurohypophysis and weakly positive in isolated cell clusters of the adenohypophysis (corresponding to acytophilic cells) (Figure 1H).

43RFa and 26RFa both promote survival and proliferation and inhibit apoptosis in INS1-E β-cells

One of the major pathological conditions, responsible for the progression of diabetes, is β-cell loss by apoptosis and decreased β-cell mass. Therefore, we tested the hypothesis as to whether 43RFa and 26RFa may protect β-cells and human pancreatic islets from the harmful effects of stimuli classically involved in diabetes, such as cytokines or glucolipotoxicity.

43RFa and 26RFa effects on survival and proliferation were assessed in INS1-E β-cells cultured in serum-free medium, in either absence or presence of the inflammatory cytokines interferon-gamma (IFN-γ), tumor necrosis factor-alpha (TNF-α) and interleukin-1 beta (IL-1β). Serum starvation
se is known to reduce survival in many cell types and cytokine synergism has been implicated in β-cell destruction and apoptosis in both type 1 and type 2 diabetes (4,25). 43RFa and 26RFa were initially tested at different concentrations (1, 10, 25, 50 and 100 nmol/L). Both peptides increased cell survival under serum starvation from 10 to 100 nmol/L, as assessed by MTT assay (Figure 2A). In cytokine-treated cells, they both promoted survival at all the concentrations tested, showing the greatest effect at 100 nmol/L (Figure 2B). Similar results were obtained by performing the Trypan blue assay (data not shown). 43RFa and 26RFa also increased cell proliferation under serum starvation at all the concentrations tested (Figure 2C), whereas with cytokines, they were effective from 25 to 100 nmol/L (Figure 2D). As for cell survival, the best proliferative effect was observed at 100 nmol/L, for both peptides, with a slight dose-response under cytokine treatment. Notably, under either serum starvation or cytokines, at 100 nmol/L the peptides increased survival and proliferation up to levels comparable to those of serum and serum-free medium, respectively.

The glucagon-like peptide 1 receptor (GLP-1R) agonist exendin-4 (Ex-4), which exerts antiapoptotic, survival and proliferative effects in β-cells (27), was used as control peptide. The survival and proliferative effects of Ex-4 were comparable to those of 43RFa and 26RFa, particularly at 100 nmol/L (Figure 2A-D).

Glucolipotoxicity, caused by excessive glucose and/or lipids, is a consequence of type 2 diabetes and a major cause of β-cell dysfunction and death (3). Based on the results obtained in cytokine-treated cells, 43RFa and 26RFa, were tested at 100 nmol/L, the concentration that was chosen for the following experiments. The peptides equally increased INS-1E β-cell survival and proliferation, that were both reduced in cells incubated with high glucose (30 mM) and palmitate (0.4 mmol/L), compared to serum-free medium alone (Figure 2E and F). These effects were similar to those of Ex-4.

As assessed by Hoechst staining of apoptotic nuclei (Figure 2G) and caspase 3 activation (Figure 2H), apoptosis was increased in cells cultured in serum free medium, compared to those in serum, and even more with cytokines or high glucose/palmitate. Both 43RFa and 26RFa, at 100 nmol/L
and similarly to Ex-4 (100 nmol/L), strongly reduced apoptosis in all experimental conditions (Figure 2G and H).

Overall, these results indicate that 43RFa and 26RFa protect pancreatic β-cells against different diabetogenic stimuli.

**43RFa and 26RFa both promote survival and inhibit apoptosis in human pancreatic islets**

Similarly to INS1-E β-cells, 43RFa and 26RFa increased survival and inhibited apoptosis in human pancreatic islets cultured for 72 h under either serum starvation, cytokine synergism or high glucose/palmitate. The peptides and Ex-4, that was used as positive control, were tested at 1, 10, 25, 50 and 100 nmol/L in islets cultured in either serum-free medium alone or with IFN-γ/TNF-α/IL-1β. Under serum starvation, 43RFa and 26RFa increased survival at 100 nmol/L, having no significant effect at lower concentrations (Figure 3A). In the presence of cytokines, they both increased cell survival from 25 to 100 nmol/L, which was the best concentration tested (Figure 3B). Conversely, Ex-4 was effective at all concentrations in both experimental conditions and in a dose-dependent manner (Figure 3A and B). At 100 nmol/L, 43RFa- and 26RFa-induced survival was similar to that of Ex-4 and a dose-response was observed under cytokine treatment (Figure 3B). 100 nmol/L was also used in islets cultured in high glucose/palmitate, where the peptides equally increased islet cell survival (Figure 3C). Apoptosis, assessed by Hoechst staining and caspase-3 activity was inhibited by both 43RFa and 26RFa, in all experimental settings and similarly to Ex-4 (Figure 3D and E).

These results indicate that, as for β-cells, in human islets 43RFa and 26RFa exert protective effects against different detrimental diabetogenic stimuli.

**43RFa survival effects involve activation of phosphoinositide 3-kinase (PI3K)/Akt and extracellular signal-regulated kinases (ERK)1/2, whereas only ERK1/2 is required for 26RFa-induced cell survival**
We next investigated the signaling pathways involved in the survival effects of the peptides, and specifically, PI3K/Akt and ERK1/2, whose activation plays a key role in β-cell growth and survival (28-30).

In INS-1E cells, 43RFa increased the phosphorylation of both Akt (at 15 min) and ERK1/2 (5 to 30 min, compared to basal time point) (Figure 4A and B). 26RFa displayed no effect on Akt, at any time point (Figure 4C), whereas it promoted ERK1/2 increase, from 5 to 60 min compared to basal (Figure 4D). Similar effects were observed in human pancreatic islets, where 43RFa induced Akt and ERK1/2 phosphorylation (both at 5 to 30 min) (Figure 4G and H), whereas 26RFa promoted ERK1/2 (5 and 15 min) (Figure 4J), but not Akt activation (Figure 4I). Accordingly, in both INS-1E and human islets cultured in serum-free medium (Figure 4E and K) or in the presence of cytokines (Figure 4F and L), preincubation with the inhibitors for either Akt (wortmannin) or ERK1/2 (PD98059) blocked the survival action of 43RFa, whereas that of 26RF was reduced only by PD98059 and not by wortmannin. These results suggest that both PI3K/Akt and ERK1/2 are involved in the survival effects of 43RFa, whereas for 26RFa, only ERK1/2 is engaged.

43RFa promotes glucose-stimulated insulin secretion and potentiates Ex-4 insulinotropic action in β-cells and human islets, whereas 26RFa inhibits insulin release and reduces Ex-4 effects

We next sought to determine 43RFa and 26RFa effects on glucose-stimulated insulin secretion (GSIS) in both INS-1E and human pancreatic islets. The peptides were tested at different glucose concentrations, either alone or in combination with Ex-4, that was used as positive control. As expected, Ex-4 increased GSIS at all glucose concentrations, in both β-cells and islets. Similarly, 43RFa increased insulin release at all glucose concentrations in β-cells and islets, to an extent comparable to that of Ex-4. 43RFa even increased basal insulin levels, without glucose stimulation. Moreover, Ex-4- and 43RFa-induced insulin secretion was further potentiated when Ex-4 and 43RFa were given together, at 15 and 25 mmol/L glucose in INS-1E cells, and at 7.5, 15 and 25
mmol/L glucose in human islets (Figure 5A and B). On the other hand, 26RFa strongly inhibited both basal and GSIS, and dramatically reduced Ex-4 insulinotropic effects at all glucose concentrations, in either INS-1E β-cells or human islets (Figure 5A and B).

To investigate the signaling pathways involved in the insulinotropic and insulinostatic effects of 43RFa and 26RFa, respectively, INS-1E β-cells and human islet cells were preincubated with either pertussis toxin (PTX), an inhibitor of Ga\(_{i/o}\) coupled receptor (31), or with NF449, a selective Ga\(_s\) protein-coupled receptor antagonist (32). 43RFa-induced increase of insulin secretion was unaffected by pretreatment with PTX, whereas it was blocked by NF449, in both β-cells and islets. Conversely, PTX restored up to control levels GSIS, that was inhibited by 26RFa, whereas NF499 had no effect on the insulinostatic action of the peptide (Figure 5C-F). These results suggest that the opposite effects of 43RFa and 26RFa on insulin secretion involve signaling through Ga\(_s\) and Ga\(_{i/o}\), respectively. Furthermore, the role of GPR103 in 43RFa and 26RFa effects on GSIS was determined by transfecting INS-1E β-cells with small interfering RNA (siRNA) targeted to GPR103. As assessed by RT-PCR, at 24 after transfection GPR103 mRNA expression was strongly reduced by siGPR103, compared with scrambled (control) siRNA (Figure 5G). The insulinotropic effect of 43RFa, assessed at both 0 and 7.5 mmol/L glucose, was completely blocked in siGPR103 cells, as compared to siControl, whereas 26RFa-induced inhibition of GSIS was unchanged in siGPR103 transfected cells (Figure 5H). These findings suggest that 43RFa promotes insulin secretion through binding to GPR103, whereas 26RFa inhibits insulin release likely through a different receptor.

43RFa, but not 26RFa, promotes cAMP increase and glucose uptake in β-cells and human pancreatic islets

cAMP elevation is linked to increased insulin secretion in pancreatic β-cells and protection from apoptosis (33,34). Therefore, we investigated the effects of 43RFa and 26RFa on cAMP levels in both INS-1E β-cells and human pancreatic islets. In INS-1E β-cells, 43RFa elevated intracellular
cAMP at 5 to 30 min with respect to basal (time 0), but not at 60 min, where cAMP returned to basal levels (Figure 6A). Conversely, 26RFa reduced cAMP levels, at the same time points (Figure 6B). Similarly, in human pancreatic islets, cAMP was increased by 43RFa at 15 and 30 min and inhibited by 26RFa, at 5 to 30 min (Figure 6C and D). The cAMP-increasing agent forskolin was used as positive control.

Glucose uptake by pancreatic β-cells, through the glucose transporter GLUT2 and glucokinase activity, is essential for the normal insulin secretory response to hyperglycemia (35,36). To further ascertain the stimulatory and inhibitory roles of 43RFa and 26RFa, respectively, on GSIS, glucose uptake was determined in INS-1E β-cells and human pancreatic islets. In INS-1E β-cells, at 10 min incubation 43RFa increased glucose 2-[3H]-deoxyglucose uptake at 10 nmol/L and showed a remarkable effect at 100 nmol/L, similarly to Ex-4 and to the cAMP analog 8-Br-cAMP. At 5 min, 43RFa had no effect at 10 nmol/L, but stimulated glucose uptake at 100 nmol/L. Conversely, 26RFa had no effect, at any concentration and incubation time tested (Figure 6E). Similarly, in human pancreatic islets, that were tested for 10 min with 100 nmol/L of each peptide, 43RFa, but not 26RFa, increased 2-deoxyglucose uptake (Figure 6F).
Discussion

The present study shows that the RFamide peptides 43RFa and 26RFa both display survival and antiapoptotic effects in pancreatic β-cells and human pancreatic islets. Furthermore, 43RFa stimulated, whereas 26RFa inhibited the insulin response to glucose and Ex-4, in both β-cells and human pancreatic islets. These effects involved activation of distinct G proteins and different regulation of cAMP levels and glucose uptake.

In mammals, the mRNAs encoding for the 43RFa and 26RFa precursor (QRFP) and for its receptor GPR103, are mainly expressed in the brain, particularly in the hypothalamus, and in different peripheral organs (6,9). QRFP mRNA has been found expressed in mouse and rat pancreas, but not in human pancreas (37). GPR103 was reported to be absent in human pancreas and expressed at very low level in mouse pancreas (6,7,37). Here we show QRFP and GPR103 expression in both rat INS-1E β-cells and human pancreatic islets, either at mRNA or protein level. Interestingly, in human islets, GPR103 protein co-localized with insulin, suggesting direct interaction of 43RFa and 26RFa with their receptor in β-cells. Furthermore, QRFP expression in β-cells and human islets may suggest autocrine/paracrine effects of the peptides. Accordingly, 26RFa autocrine/paracrine effects have been recently proposed in prostate cancer cells, where the peptide induced cell migration and neuroendocrine differentiation, and in macrophages and adipocytes, where both 43RFa and 26RFa promoted adipogenesis (10,11,22). Future studies will help elucidate whether autocrine/paracrine mechanisms may also occur in pancreatic islets.

We firstly report here the survival and antiapoptotic effects of 43RFa and 26RFa. Pancreatic β-cells and human pancreatic islets were chosen as in vitro models because, besides the survival actions, we were interested in the metabolic effects of these peptides. Indeed, a role in adipogenesis, lipid metabolism and inflammation, as well as an association between GPR103 and QRFP expression and obesity, have been recently described (10,11). Moreover, as for QRFP, other orexigenic peptides such as ghrelin, exert antiapoptotic and metabolic effects in different cell types, including pancreatic β-cells and human islets (24,38,39). Here, both 43RFa and 26RFa increased survival and
inhibited apoptosis of INS-1E β-cells and human pancreatic islets. They also increased proliferation of INS-1E β-cells, that was not investigated in human islets, due to the low proliferative rate of human β-cells in vitro (40,41). Besides serum starvation, which induces β-cell death (24,42), β-cells and human islets were challenged with diabetogenic stimuli, such as cytokine synergism and glucolipotoxicity, which are major causes of β-cell dysfunction and death in both type 1 and 2 diabetes (1-4). Interestingly, 43RFa and 26RFa both displayed proliferative, survival and antiapoptotic actions, comparable to those of the long acting GLP-1 agonist Ex-4 (27).

Phosphorylation of either PI3K/Akt or ERK1/2 has been shown to mediate β-cell proliferation and survival in response to different external signals, including incretin hormones (24,27,30,43). Here, 43RFa and 26RFa, although displaying similar survival effect in β-cells and human islets, differently regulated these signaling pathways. In fact, whereas 43RFa increased the phosphorylation of both PI3K/Akt and ERK1/2, 26RFa increased ERK1/2, but not PI3K/Akt. Accordingly, inhibition of PI3K and ERK1/2 phosphorylation with wortmannin and PD98059, respectively, blocked 43RFa-induced survival, whereas only ERK1/2 inhibitor decreased 26RFa-induced survival. These results indicate that ERK1/2, and not PI3K/Akt, is required for 26RFa survival effect, whereas 43RFa uses both pathways. Notably, a cross talk between ERK1/2 and cAMP in pancreatic β-cell survival has been previously described (29). However, here only 43RFa elevated cAMP levels, suggesting that 26RFa-induced activation of ERK1/2 is independent of the cAMP pathway.

To further understand the effects of 43RF and 26RFa in β-cells we next determined their role on insulin secretion. To our knowledge, 43RFa effect on insulin secretion has never been studied. Instead, 26RFa was recently shown to inhibit glucose- as well as arginine- and Ex-4-induced insulin secretion in perfused rat pancreas (23). Here, we firstly show that 43RFa promoted insulin secretion in INS-1E β-cells and human islets, both in basal conditions and in the presence of either different glucose concentrations or Ex-4. Conversely, 26RFa showed opposite effects and inhibited both glucose- and Ex-4-induced insulin secretion, in agreement with the previous findings (23). 26RFa
was used here at 100 nmol/L, which was also the best concentration to promote survival and prevent apoptosis. Conversely, 10 nmol/L was the concentration used in the study by Egido et al., where the experiments were performed in perfused rat pancreas, at variance with our in vitro models (23).

Interestingly, either 43RFa or Ex-4 alone, showed comparable insulinotropic action in both β-cells and human islets. Moreover, insulin secretion was markedly increased when 43RFa and Ex-4 were administered together, particularly at the highest concentrations of glucose. On the other hand, besides exerting inhibitory action per se, 26RFa hampered Ex-4 insulinotropic effect. These results imply that 43RFa may either potentiate Ex-4 action or act synergistically with Ex-4 to increase insulin secretion, whereas 26RFa behaves in an opposite manner. They also suggest clinical implications in diabetes, as 43RFa may become a therapeutic target for enhancing β-cell function, alone or in combination with incretin hormones, whereas inhibition of 26RFa effect may prevent loss of β-cell competence.

Activation of G protein-coupled receptors (GPCRs) by metabolites and hormones can modulate glucose-induced insulin secretion through different pathways. Ga\textsubscript{s} coupled receptors increase insulin secretion through stimulation of adenylyl cyclase and activation of protein kinase A (PKA) by cAMP. In turn, Ga\textsubscript{i/o} coupled receptors reduce insulin release by inhibition of adenylyl cyclase and cAMP formation and also through inhibition of voltage-dependent Ca\textsuperscript{2+} channels involving the βγ subunits of Ga\textsubscript{i/o} (27,44). We found here that the Ga\textsubscript{s} antagonist NF449 blocked the insulinotropic effect of 43RFa, but not the insulinostatic action of 26RFa. Conversely, the Ga\textsubscript{i/o} inhibitor PTX did not affect 43RFa-induced insulin release, but blocked the inhibitory effect of 26RFa, in both β-cells and human islets. This in agreement with the findings by Egido et al. (23), except for the fact that here PTX was used in combination with glucose only, and not with Ex-4. Of note, PTX alone had no effect here on GSIS, whereas with 26RFa, insulin levels were found increased. This is likely because glucose alone, in the absence of GPCR ligands, causes inhibition of ATP-regulated K\textsuperscript{+} channels, opening of voltage-dependent Ca\textsuperscript{2+} channels and increase in
intracellular Ca^{2+} concentration ([Ca^{2+}]_i), which triggers exocytosis of insulin-containing vesicles (45). Conversely, in the presence of 26RFa, PTX may restore cAMP levels, which are blunted by the peptide, as well as ([Ca^{2+}]_i), and may thus contribute to increase insulin release.

Our results indicate that the opposite effects of 43RFa and 26RFa on insulin secretion are due to activation of distinct pathways. In fact, 43RFa promotes insulin secretion, at least in part, through G_{as} and adenylyl cyclase/cAMP increase, whereas 26RFa inhibits insulin release through G_{ai/o} and inhibition of cAMP formation. This assumption is further sustained by our results showing that 43RFa increased, whereas 26RFa inhibited intracellular cAMP levels in both INS-1E β-cells and human pancreatic islets. At variance with these findings, 43RFa has been recently found to decrease forskolin-induced cAMP in human adrenocortical cells, likely through G_{q-mediated mechanisms (20). 26RFa, instead, in line with our results, was shown to decrease cAMP production in Chinese hamster ovary (CHO)- or human embryonic kidney (HEK293)-GPR103 transfected cells, suggesting GPR103 coupling to G_{i/o} and G_{q} (7). In addition, we found here that similarly to the cAMP analog 8-Br-cAMP and to Ex-4, 43RFa, but not 26RFa, increased glucose uptake, which is required in the initial step of GSIS by pancreatic β-cells and has been shown to be impaired in both type 1 and type 2 diabetes (35,46). Collectively, these findings further support the positive role of 43RFa on β-cell function, which is opposed to that of 26RFa.

An open question during the study was why two peptides, that likely bind the same receptor, exert opposite effects on insulin secretion. To solve this issue, we knocked down GPR103 expression in INS-1E β-cells using specific siRNA, and found complete inhibition of 43RFa insulinotropic action, but no effect on 26RFa-induced insulin inhibition. This finding suggests that in β-cells, only 43RFa signals through GPR103, whereas 26RFa interacts with a different receptor. Accordingly, 26RFa was found to display moderate affinity and selectivity for NPFF2, the receptor of neuropeptide FF, another member of the RFamide family (47). Moreover, NPFF and prolactin-releasing peptide (PrRP), both with high affinity for NPFF2 receptor (9), displaced [^{125}I]-26RFa binding in brain regions enriched with NPFF2 receptor (48). Interestingly, NPFF2 receptor is coupled to G_{i/o} protein.
in agreement with our finding showing loss of 26RFa insulinostatic effect in β-cells and human islets treated with PTX.

In conclusion, this study shows a new role for 43RFa and 26RFa in promoting β-cell and human islet cell survival, while exerting opposite effects on insulin secretion, via either \( \text{G} \alpha_s/\text{cAMP} \) elevation or \( \text{G}_i/\text{cAMP} \) inhibition, respectively. The effects on insulin secretion require GPR103 binding for 43RFa, but not for 26RFa. These findings provide new perspectives and comprehension on the role of these peptides, whose functions, particularly at the peripheral level, are still quite unknown. In addition, 43RFa and 26RFa may become novel therapeutic targets for increasing β-cell mass and function in metabolic diseases such as diabetes and obesity.
Acknowledgements

This study was supported by the University of Torino (Ex-60% 2008, to R.G.), by the Ministero dell’Istruzione, Università e della Ricerca Scientifica e Tecnologica [Italian Ministry of Instruction and Research (MIUR: PRIN 2008EFHJ5H_02 to R.G.; PRIN 2010B5B2NL to E.G.)]; by Compagnia di San Paolo (2007 to E.G and 2011 to R.G.), by Sanofi-aventis S.p.A., Italy and by Studio delle Malattie Endocrino Metaboliche (SMEF) Foundation, Turin, Italy. Human islets were provided through the JDRF Award 31-2008-416 (ECIT; Islet for Research Program).

R.G. contributed to study design and interpretation and wrote the manuscript; F.S., L.T., D.G. I.G., L.B. researched data and reviewed/edited the manuscript; R.N and L.P. provided the human islets and reviewed/edited the manuscript; M.P.G. and G.A. researched data on Ca^{2+} transients and reviewed/edited the manuscript; M.V. and M.P. contributed with the immunohistochemical studies and reviewed/edited the manuscript; J.L., H.V., H.O. and E.G. critically revised the paper and helped with the Discussion. R.G is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. No potential conflicts of interest relevant to this article were reported. The authors are grateful to Eleonora Gargantini, Jessica Giorcelli, Cristina Grande, Ida Rapa and Marina Taliano (all from the University of Torino) for their technical support and to the Neuroscience Institute of Turin (NIT) and the Italian Group of Neuroendocrine Sciences (GISNe).
Figure legends

Figure 1. Gene and protein expression of GPR103 and QRFP (the RF43a and 26RFa precursor) in INS1-E β-cells and human pancreatic islets (H. islets). (A, B) GPR103 (A) and QRFP (B) mRNA assessed by RT-PCR. Rat brain (A and B) and adult hippocampal progenitor cells (AHPs) (A) were used as positive control; 18s rRNA was used as internal control. (C and D) GPR103 (C) and QRFP (D) protein expression assessed by Western blot on whole lysates from INS-1E β-cells or human pancreatic islets. Equal protein loading was determined by reprobing with antibodies to β-actin. The blots are representative of three independent experiments. (E) Immunofluorescent staining of GPR103 in INS-1E β-cells. GPR103 is shown in red and nuclei in blue (DAPI); AHPs were used as positive control. Each image is representative of three independent experiments (X40). (F) GPR103 expression in human pancreatic islets assessed by immunofluorescent staining. Insulin is shown in green and GPR103 in red (X40) (n=3). Merge shows colocalization of GPR103 and insulin (yellow), as indicated by arrows. (G) Immunohistochemical analysis of GPR103 and QRFP expression in INS-1E β-cells. Cells incubated without the primary antibody were used as negative control (-). (H) Immunohistochemical analysis of GPR103 and QRFP expression in human pancreas; pituitary was used as positive control (n=3).

Figure 2. 43RFa and 26RFa effects on survival, proliferation and apoptosis of INS-1E β-cells. The cells were cultured in serum-free medium (SF) for 12 h, then incubated for a further 24 h without or with the cytokines (CK) TNF-α/IFN-γ/IL-1β (100, 50 and 5 ng/ml respectively), or with high glucose (30 mmol/L) and palmitate (0.4 mmol/L) (P/G), and either with 43RFa, 26RFa or Ex-4. The peptides were added 40 min prior to CK or P/G. (A-D) Cell survival and proliferation (assessed by MTT and BrdU, respectively) in the absence (A, C) or presence (B, D) of CK and the peptides, at the indicated concentrations. Data are expressed as percent of control (SF) and are the mean ± SE of five replicates (n=3) (*P<0.05, **P<0.01 vs. SF; #P<0.05, ##P<0.01 vs. CK; ns, not significant). Cell survival (E) and cell proliferation (F) in cells cultured in SF medium, alone or with P/G and the
indicated peptides (100 nmol/L each). Results are expressed as percent of control (SF); n=3. (**P<0.01 vs. SF; ##P<0.01 vs. P/G). (G) Apoptosis assessed by counting condensed/fragmented Hoechst stained apoptotic nuclei. Values are expressed as percent of apoptotic cells and are the mean ± SE of duplicate determinations (500 cells each) from three independent experiments. (H) Apoptosis assessed as caspase-3 activation and expressed as percent of control (SF). (For both G and H: **P<0.01; ##P<0.01 vs. SF; n=3). Each peptide was used at 100 nmol/L.

**Figure 3.** 43RFa and 26RFa effects on survival and apoptosis of human pancreatic islets. Islet cells were incubated for 72 h in the presence of serum or in serum-free medium (SF), either alone or with IFN-γ/TNF-α/IL-1β (CK) (5 ng/ml each) and 43RFa or 26RFa. (A, B) Cell viability assessed by MTT in islets cultured in serum-free medium (SF), alone (A) or with cytokines (CK) (B) and the peptides at the indicated concentrations. Results are expressed as percent of control (SF, n=3) (*P<0.05, **P<0.01 vs. SF for A or vs. CK for B; ns, not significant). (C) Cell survival (MTT) in human islets treated in SF medium alone or with high glucose (30 mmol/L)/palmitate (G/P) (0.4 mmol/L) and the indicated peptides (100 nmol/L each). Results are expressed as percent of control (SF); n=3. (*P<0.05, **P<0.01 vs. SF; ##P<0.01 vs. P/G). (D) Apoptosis assessed by Hoechst staining. Values are expressed as percent of apoptotic cells from duplicate determinations (500 cells each, n=3). (E) Apoptosis assessed by caspase-3 activation. Results are expressed as percent of control (SF). (For both D and E: *P<0.05, **P<0.01; ##P<0.01 vs. SF; n=3). Each peptide was used at 100 nmol/L.

**Figure 4.** 43RFa and 26RFa effects on activation of PI3K/Akt and ERK1/2. Akt and ERK1/2 phosphorylation was assessed by Western blot on whole lysates from INS-1E cells or human pancreatic islets stimulated with either 100 nmol/L 43RFa (A-B and G-H) or 100 nmol/L 26RFa (C-D and I-J), for the indicated times (top panels). Equal protein loading was determined by reprobing
with antibodies to the respective total proteins (*bottom panels*). Blots are representative of three independent experiments. Graphs show the densitometric analysis of phosphorylated proteins normalized to total proteins and reported as percent of basal (*P<0.05, **P<0.01; ns, not significant). (E and F) Cell survival assessed by MTT in INS-1E cells that were starved for 12 h, then incubated for a further 24 h in the absence or presence of either 43RFa (43) or 26RFa (26) (100 nmol/L each), and either without (serum-free medium, SF) (E) or with TNF-α/IFN-γ/IL-1β (cytokines, CK) (100, 50 and 5 ng/ml respectively) (F), and wortmannin (WM) (100 nmol/L) or PD98059 (40 µmol/L). Inhibitors were added 30 and 60 min before the peptides and cytokines, respectively. **P<0.01 vs. SF (E) or vs. CK (F); #P<0.01 vs. 43; §P<0.05 vs. 26; ns, not significant; n=3. (K and L) Cell survival assessed by MTT in human pancreatic islets incubated for 72 h in the presence of serum or in serum-free medium (SF), either alone (K) or with IFN-γ/TNF-α/IL-1β (CK) (5 ng/ml each) and 43RFa or 26RFa (100 nmol/L each). Inhibitors were added as for E and F. **P<0.01 vs. SF (K) or vs. CK (L); #P<0.05, #P<0.01 vs. 43; §P<0.05 §§P<0.01 vs. 26; ns, not significant; n=3.

**Figure 5.** 43RFa and 26RFa regulate insulin secretion in INS-1E β-cells and human pancreatic islets. Insulin release was assessed by RIA in conditioned medium from INS-1E β-cells or islets, incubated alone or with 2 mmol/L glucose for 1 h and then for a further 1 h with the indicated concentrations of glucose, in the presence or absence of either 43RFa, 26RFa, Ex-4 or a combination of Ex-4/43RFa or Ex-4/26RFa (100 nmol/L each). (A) Insulin secretion in INS-1E β-cells. (B) Insulin secretion in human pancreatic islets. For A and B, values are the means ± SE of triplicate determinations from at least three independent experiments (*P<0.05, **P<0.01 vs. Control at each glucose concentration; #P<0.05, #P<0.01). (C-F) Insulin secretion in INS-1E cells (C and D) and human pancreatic islets (E and F) in either absence or presence of pertussis toxin (PTX) (50 ng/ml) or NF449 (10 µmol/L). *P<0.05, **P<0.01 vs. control (c) at each glucose concentration; #P<0.05, #P<0.01; ns, not significant. (G) GPR103 mRNA assessed by RT-PCR in
INS-1E β-cells after 24 h transfection with either nonsilencing siRNA (siControl) or siRNA to GPR103 (siGPR103). As expected, the amplified products corresponded to 318 bp for rat GPR103, and 199 bp for 18s rRNA, that was used as internal control. (H) Insulin secretion in siControl- or siGPR103-transfected INS-1E β-cells, incubated for 1 h with the indicated concentrations of glucose, and with 43RFa or 26RFa (100 nmol/L each). Values are the means ± SE of triplicate determinations from at least three independent experiments (*P<0.05 vs. Control at each glucose concentration; #P<0.05).

**Figure 6.** 43RFa and 26RFa effects on intracellular cAMP and glucose uptake. (A-D) Serum-starved INS-1E cells and human islets were cultured for the indicated times with either 100 nmol/L 43RFa or 100 nmol/L 26RFa, in the presence of the phosphodiesterase inhibitor IBMX (100 µmol/L), that was added 30 min before stimulation. Forskolin (Forsk) (50 µmol/L for 1 min) was used as positive control. Results are the mean ±SE of three independent experiments performed in triplicate (*P<0.05, **P<0.005 vs. basal time point; ns, not significant). (E) 2-[3H]-deoxyglucose glucose uptake assessed in serum starved INS-1E cells cultured for 5 or 10 min with either Ex-4, 43RFa or 26RFa, at the indicated concentrations. (F) 2-[3H]-deoxyglucose glucose in human pancreatic islets cultured for 10 min with the peptides, that were used at 100 nmol/L each. 8-Br-cAMP was used at 1 mmol/L. For E and F: n=3; *P<0.05**P<0.005, ***P<0.001; ns, not significant vs. control.
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Figure 1

A
GPR103
100 rRNA

B
QRFP
100 rRNA

C
GPR103
p-actin

D
QRFP
p-actin

E
DAPI
GPR103
Merge

F
Human pancreatic islets

G
Control

H
Human pituitary
Human pancreas

165x272mm (150 x 150 DPI)
Figure 2

INS-1E

A. Serum starvation

B. Cytokines

C. Serum starvation

D. Cytokines

E. SF, PG

F. SF, PG

G. Apoptosis (%)

H. Caspase-3 activity (% of control)

260x430mm (300 x 300 DPI)
Figure 4

INS-1E

A

43RFa

B

26RFa

C

P-Akt

D

ERK1/2

P-Akt

Akt

ERK1/2

P-Akt

Akt

ERK1/2

E

serum starvation

F

cytokines

Human pancreatic islets

G

43RFa

H

26RFa

I

P-Akt

J

ERK1/2

P-Akt

Akt

ERK1/2

P-Akt

Akt

ERK1/2

K

serum starvation

L

cytokines

Cell survival (% of control)

183x240mm (300 x 300 DPI)
Figure 5

A

INS-1E

B

Human pancreatic islets

C

INS-1E

D

INS-1E

E

Human pancreatic islets

F

Human pancreatic islets

G

H

GPR103

18s rRNA

siControl

siGPR103

242x317mm (300 x 300 DPI)