Antineoplastic effects of rosiglitazone and PPARγ transactivation in neuroblastoma cells

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Neuroblastoma (NB) is the most common extracranial solid tumour in infants. Unfortunately, most children present with advanced disease and have a poor prognosis. In the present study, we evaluated the role of the peroxisome proliferator-activated receptor γ (PPARγ) agonist rosiglitazone (RGZ) in two NB cell lines (SK-N-AS and SH-SY5Y), which express PPARγ. Rosiglitazone decreased cell proliferation and viability to a greater extent in SK-N-AS than in SH-SY5Y. Furthermore, 20 μM RGZ significantly inhibited cell adhesion, invasiveness and apoptosis in SK-N-AS, but not in SH-SY5Y. Because of the different response of SK-N-AS and SH-SY5Y cells to RGZ, the function of PPARγ as a transcriptional activator was assessed. Noticeably, transient transcription experiments with a PPARγ responsive element showed that RGZ induced a three-fold increase of the reporter activity in SK-N-AS, whereas no effect was observed in SH-SY5Y. The different PPARγ activity may be likely due to the markedly lower amount of phosphorylated (i.e. inactive) protein observed in SK-N-AS. To our knowledge, this is the first demonstration that the differential response of NB cells to RGZ may be related to differences in PPARγ transactivation. This finding indicates that PPARγ activity may be useful to select those patients, for whom PPARγ agonists may have a beneficial therapeutic effect.

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Thiazolidinediones (TZDs) are a class of molecules, which activate the nuclear receptor peroxisome proliferator-activated receptor γ (PPARγ) (Desvergne and Wahli, 1999) and promote association with the 9-cis retinoic X receptor (RXR) to form functional heterodimers that recognise its cognate DNA response element within target genes (Jude-Aubry et al, 1997; Reginato et al, 1998). In addition to their well-known effects on glucose homeostasis (Yki-Järvinen, 2004), TZDs have been shown to have anti-inflammatory (Jiang et al, 1998, Marx et al, 2000) and antineoplastic effects (Koeffler, 2003, Grommes et al, 2004). The latter effect is in agreement with the demonstration that PPARγ/RXR signalling exerts an important role in inhibiting cell proliferation and/or in inducing apoptosis (Grommes et al, 2004). In addition, loss-of-function mutations of the PPARγ gene have been found in human colon and thyroid cancer (Sarraf et al, 1999; Kroll et al, 2000). Therefore, PPARγ has been regarded as a target for anticancer therapy and clinical trials with TZDs for the treatment of human malignancies involving different organs and tissues, such as the prostate (Hisatake et al, 2000; Mueller et al, 2000), the colon (Kulke et al, 2002), the breast (Burstein et al, 2003) and the adipose tissue (Demetri et al, 1999; Debrock et al, 2003) have been initiated in recent years. Peroxisome proliferator-activated receptor γ is expressed in tumours of the nervous system, such as astrocytomas, glioblastomas and neuroblastomas (Han et al, 2001; Nwako and Robbins, 2001; Strakova et al, 2004). Neuroblastoma (NB) is the most common extracranial solid tumour in children and it has a heterogeneous clinical presentation and course (Shimada and Joshi, 1997). Unfortunately, most children with NB present with advanced disease. More than 60% of patients with high-risk features will have a poor prognosis despite intensive therapy. Thus, research efforts to understand the biologic basis of NB and to identify new and more effective therapies are essential to improve the outcome for these children. Retinoids have been shown for instance to interfere with cell growth and to induce apoptosis in NB cells (Melinò et al, 1997, Voigt and Zintl, 2003) and preliminary clinical trials with retinoids in NB resulted in improved event-free survival in high-risk patients, with limited toxic effects (Garaventa et al, 2003; Reynolds et al, 2003).

The aim of our study was to evaluate the role of the PPARγ agonist rosiglitazone (RGZ) on cell growth, adhesion, invasiveness and apoptosis in two different NB cell lines (SK-N-AS and SH-SY5Y).
SH-SY5Y), which express PPARγ (Servidei et al., 2004, Valentina et al., 2005). Furthermore, transient transfection experiments with a peroxisome proliferator response element-luciferase reporter plasmid were performed, in order to determine whether cell response to RGZ was related to the level of PPARγ transcriptional activity. Finally, gene sequencing of the PPARγ gene was performed and the amount of expression of PPARγ (total and phosphorylated) was evaluated and correlated to its transcriptional activity in the two NB cell lines.

MATERIALS AND METHODS

Materials

Human NB cell lines SH-SY5Y and SK-N-AS (American Type Culture Collection, Manassas, VA, USA) were made available by the laboratory of Dr CJ Thiele. All the reagents for cell cultures were from Sigma Chemical Co. (St Louis, MO, USA). Tissue plastic-ware was from Bibby Sterilin (Staffordshire, UK). Rosiglitazone was purchased from Alexis (San Diego, CA, USA).

Cell proliferation

DNA synthesis in NB cells was evaluated by [3H]-thymidine incorporation assay. The cells were seeded in 24-well plates and treated with RGZ for 24 and 48 h. During the last 4 h, the cells were pulsed with 1 μCi well–1 [3H]-thymidine (Amersham Biosciences, Little Chalfont, Buckinghamshire, UK). Finally, the cells were washed with PBS and fixed by cold trichloroacetic acid (5% in distilled water). A 600 μl well–1 portion of extraction buffer (NaOH 0.25 N/SDS 0.1%) was added for 30 min at 37°C for DNA extraction. Successively, after pH neutralisation with 6 N HCl, the samples were analysed by a β-counter (Betamatic V, Kontron Instruments Ltd, Betchley, UK). The PPARγ antagonist 2,2-bis[4-(glicycoloxy) phenyl]propane (BADGE, Sigma, Milan, Italy) was used at the concentration of 20 μM and it was added to the cultured cells 30 min before RGZ exposure.

For cell proliferation assay, the cells were seeded in six-well plates. After 24 h, the cells were washed with PBS and different doses of RGZ were added in RPMI medium 1% FBS for 24 or 48 h. Then, the cells were detached (trypsin 0.25%/EDTA) and the cell number was determined by a haemocytometer.

Cell viability

Cell viability following exposure to different RGZ doses (0.1–20 μM) was determined by MTS assay (Promega Corporation, Madison, WI, USA), as described previously (Benvenuti et al., 2005). Briefly, the cells were seeded in 96-well plates and exposed to RGZ for 48 h. The assay was then performed according to the manufacturer’s instructions. The samples were analysed by an ELISA plate reader (Seac-Radim, Moncalieri, Italy) at 490 nm wavelength. The PPARγ antagonist BADGE (Sigma) was used at the concentration of 20 μM and it was added to the cultured cells 30 min before RGZ exposure.

Cell adhesion assay

The cells were plated in 96-well plates and different doses of RGZ (0.1–20 μM) were added for 24 and 48 h. After washing with PBS, 100 μl of Bengal Rose stain (0.25% in PBS) was added to each well for 5 min at room temperature (Gamble and Vadas, 1988). After aspiration of the stain and washing with PBS, the stain was released by adding 200 μl well–1 of an ethanol/PBS (1:1) solution for 30 min. The samples were analysed by an ELISA plate reader (Seac-Radim) at 570 nm wavelength.

Invasion assay

Cell invasion was evaluated using the Chemicon Cell Invasion Assay (Temecula, CA, USA). The assay is performed in a 24-well tissue culture plate with 12 cell culture inserts and each Invasion Chamber insert contains an 8 μm pore size membrane over which a thin layer of extracellular matrix is dried. The cells were suspended in serum-free medium and RGZ (20 μM) was added. Successively, after rehydration of the insert, the cells were plated in each Invasion Chamber insert. After 24 h, the noninvading cells on the upper surface of the filter were removed, whereas the invasive cells were stained with the Cell Stain Solution provided with the kit, eluted with 150 μl of 1% acetic acid, transferred onto a microplate and analysed by an ELISA plate reader (Seac-Radim) at 540 nm wavelength.

Quantitation of matrix metalloproteinase-9 (MMP-9) and tissue inhibitor of matrix metalloproteinase-1 (TIMP-1) mRNA

The measurement of MMP-9 and TIMP-1 mRNA was performed by using a multiplex quantitative real-time RT–PCR method, based on TaqMan technology, as described previously (Cioppi et al., 2004). Standard curves for MMP-9 and TIMP-1 mRNAs consisted of serial 1:10 dilutions from 2.5 × 102 to 2.5 × 106 fg total RNA from the fibrosarcoma cell line HT1080 (American Type Culture Collection).

ELISA assay for MMP-9

Matrix metalloproteinase-9 was measured in conditioned medium using an ELISA-based commercial kit (Pharmacia, Uppsala, Sweden), following the manufacturer’s instructions. Matrix metalloproteinase-9 can be measured in a range of 1–32 ng ml–1, and the sensitivity of the assay is 0.6 ng ml–1.

Immunocytochemistry and flow cytometry analysis for cleaved caspase 3

The amount of caspase 3 immunoreactive cells was determined as described previously (Benvenuti et al., 2005). Briefly, the cells were seeded in chamber slides and exposed to RGZ (20 μM). After 48 h, the cells were fixed in paraformaldehyde, then incubated with a rabbit polyclonal antibody against human cleaved caspase 3 (Asp 175) (Cell Signalling Technology Inc., Beverly, MA, USA) and subsequently with a biotinylated secondary antibody. The reaction product was visualised by ABC peroxidase-based detection protocol and AEC kit (Vectastain ABC kit, Vector Laboratories Inc., Burlingame, CA, USA). Finally, the cells were counterstained with haematoxylin and apoptotic cells/field were counted in 10 fields (×40).

Moreover, cleaved caspase 3 was evaluated by flow cytometry, which was performed as described previously (Carloni et al., 1998). Briefly, the cells were seeded onto 100 mm Petri dishes at the density of 3 × 106 cells and incubated for 24 h in complete medium. Then, RGZ (20 μM) was added and the cells were incubated at 37°C in humidified atmosphere (5% CO2/95% air). After 48 h, the cells were detached (trypsin 0.25%/EDTA) and fixed with 1% paraformaldehyde in distilled H2O for 10 min at 37°C. After permeabilisation in ice-cold 90% methanol, the cells were incubated with a rabbit polyclonal antibody against cleaved caspase 3 (Asp 175) (Cell Signalling Technology) or with a nonspecific IgG as control at 1:25. Subsequently, the cells were washed with incubation buffer and incubated with a FITC-conjugated goat anti-rabbit secondary antibody (1:100) (Southern Biotechnology Associates Inc., Birmingham, AL, USA) for 30 min. FITC green fluorescence was detected at 515–555 nm using a FL-1 detector of a FACScan.
flow cytometer (Becton Dickinson, Mountain View, CA, USA) equipped with a 15 mW argon-ion laser for excitation. Debris were gated out by establishing a region around the population of interest on the Forward Scatter vs Side Scatter dot plot. For each sample, 10,000 events in the region of interest were recorded at a flow rate of 200–300 cells/s\(^{-1}\). Data were processed with analysis software LYSYS II (Becton Dickinson).

**Transient transfection of NB cells**

In order to determine the activity of PPAR\(\gamma\) in NB cells, the cells were transfected at the density of 4 \(\times\) 10\(^6\) cells (SK-N-AS) and 2 \(\times\) 10\(^6\) cells (SH-SY5Y) in 100-mm dishes with 2 \(\mu\)g peroxisome proliferator response element (adipocyte response element)-\(\gamma\)_3-tk-luciferase reporter plasmid (containing three copies of the peroxisome proliferator response element from the adipocyte lipid binding protein gene ligated to a herpes simplex thymidine kinase promoter upstream from a luciferase gene), 0.15 \(\mu\)g of human PPAR\(\gamma\) expression plasmid, and 1 \(\mu\)g pSV\(_2\)CAT (a vector containing Simian virus 40 early promoter and enhancer sequences that drives a chimeric chloramphenicol acetyl transferase gene) as an internal control by calcium phosphate precipitation, as described previously (Ferruzzi et al, 2005). Twenty-four hours after transfection, the cells were treated with RGZ. Twenty-four hours later, the cells were harvested, washed twice with PBS and lysed. Fifty microlitres of cell extract was incubated with luciferase assay reagent based on the original protocol by de Wet et al (1987). The number of relative light units with a 3-s delay and a 30-s incubation was measured using a Sirius1 luminometer (Berthold Detection System, Pforzheim, Germany). Chloramphenicol acetyl transferase activity was measured as described previously (Crabb and Dixon, 1987). The conversion of chloramphenicol to its acetylated products was quantified on a \(\beta\)-scanner (Ambis System, San Diego, CA, USA).

**Sequence analysis of the PPAR\(\gamma\) gene**

Genomic DNA was extracted from cell cultures using the automated DNA extraction system Biorobot EZ1 Workstation (Qiagen, Crawley, W Sussex, UK). The entire coding region of the PPAR\(\gamma\) gene and the exon–intron boundaries were amplified by PCR, using primers sequences and PCR conditions as described previously (Costa-Guda et al, 2005). Amplicons were purified using the QIAquick purification kit (Qiagen) and direct sequencing was carried out using the Big Dye Terminator v1.1 Cycle Sequencing kit (Applied Biosystems, Warrington, Cheshire, UK).

![Figure 1](image-url)
according to the manufacturer’s instructions. Sequencing reactions were purified using the DyeEx 2.0 Spin kit (Qiagen) and samples were run on an ABI 310 capillary sequencer (Applied Biosystems).

**Western blot analysis of PPARγ expression**

The NB cells were washed and resuspended in lysis buffer. In addition, the human adenocarcinoma cell line PANC-1 and HPAC (American Type Culture Collection) were used as the positive and negative control for PPARγ expression, respectively (Galli et al., 2004). After protein measurement (Coomassie kit, Bio-Rad Laboratories, Hercules, CA, USA), 30 µg of protein was diluted in 2 x Laemmli’s reducing sample buffer, incubated at 95 °C for 5 min and loaded onto a 10% polyacrylamide-bisacrylamide gel. After SDS–PAGE, proteins were electroblotted into nitrocellulose (Sigma Chemical Co.). Equivalent protein loading was verified by staining parallel gels with Coomassie R. After blocking in 5% skimmed milk for 2 h, nitrocellulose membranes were washed and then immunostained with a rabbit anti-human PPARγ antibody (1 : 1000) (Santa Cruz Biotechnology, Santa Cruz, CA, USA) followed by a secondary anti-mouse IgG antibody (1 : 2000) (New England Biolabs, Beverly, MA, USA). The antibody-reacted proteins were revealed by LumiGLO chemiluminescent reagent and peroxide (New England Biolabs).

**PPARγ phosphorylation assay**

Protein extraction was performed by boiling the cell pellet in 10% glycerol, 2% SDS and 50 mM Tris-HCl (pH 7.5) for 10 min. The lysate was diluted 1 : 10 with RIPA-like buffer containing 50 mM Tris-HCl (pH 7.5), 1% Triton X-100, 0.5% sodium deoxycholate, 150 mM NaCl, 5 mM EDTA, 25 mM β-glycerophosphate, 1 mM Na3VO4, 100 µM PMSF, supplemented with Complete Protease Inhibitor Cocktail (Roche Applied Science, Milan, Italy). SDS was then added to achieve a final concentration of 1% and immunoprecipitation was performed with a rabbit anti-human PPARγ antibody (1 : 100) (Santa Cruz Biotechnology) at 4 °C, for 3 h. Immunocomplexes were recovered by incubation with Protein-A Sepharose (Sigma) for additional 16 h at 4 °C. The immunocomplexes were then dissociated by boiling for 5 min in Laemmli’s buffer, the beads were collected by centrifugation and SDS–PAGE was performed with the supernatant. The proteins were electroblotted to nitrocellulose transfer membrane Protean (Whatman, VWR International, Milan, Italy) and were detected by incubating the filter with the anti-PPAR-γ antibody or with an anti-phosphoserine mouse monoclonal antibody (1 : 400) (clone PSR-45, Sigma) followed by a secondary anti-rabbit IgG (1 : 2000) or anti-mouse IgG (1 : 2000) antibody, respectively (Amersham Biosciences). Detection of the protein bands was performed using the Amersham ECL plus Kit (Amersham Biosciences).

Figure 2  (A and B) Role of RGZ on SK-N-AS (A) and SH-SY5Y (B) cell viability, as assessed by MTS assay (see Materials and Methods). *P < 0.05 vs control untreated cells (C). (C and D) Effect of different concentrations and time of exposure to RGZ (24 or 48 h) on SK-N-AS (C) and SH-SY5Y (D) cell adhesion, as assessed by Bengal Rose assay (see Materials and Methods). *P < 0.05 vs control untreated cells (C).
RESULTS

Effects of RGZ treatment on cell proliferation

The effects of RGZ on ³[H]-thymidine incorporation in the two NB cell lines SK-N-AS and SH-SY5Y are reported in Figure 1. In SK-N-AS (Figure 1A), a significant reduction of DNA synthesis was determined by exposure to RGZ (1–20 μM) for 24 and 48 h, whereas in SH-SY5Y (Figure 1B), this effect was observed only after a 48-h exposure.

With regard to cell counts, a significant antiproliferative effect was observed following a 24- or 48-h treatment with RGZ (10 and 20 μM) in SK-N-AS (Figure 1C). In SH-SY5Y, a significant reduction of the cell number was observed only after a 48-h treatment with 20 μM RGZ (Figure 1D), although to a lower extent than in SK-N-AS. Only a higher concentration of RGZ (80 μM) was able to determine in SH-SY5Y a reduction of the cell number (54.4 ± 1.3%, mean ± s.e. vs untreated cells, considered as 100%) similar to the decrease determined by the same dose of RZG in SK-N-AS (57.9 ± 1.4%) (data not shown).

Effects of RGZ on cell viability and adhesion

The effect of RGZ on NB cell viability is shown in Figure 2A and B. In SK-N-AS, cell viability was significantly reduced after a 48-h exposure to RGZ (0.1–20 μM) and 20 μM RGZ determined the maximal inhibition (54%) (Figure 2A). In SH-SY5Y cells, RGZ (0.1–20 μM) was also able to reduce cell viability, but the decrease was limited to 13% (20 μM RGZ) (Figure 2B).

Moreover, the effect of RGZ on cell adhesion was determined. Interestingly, in SK-N-AS cells RGZ significantly induced an inhibition of cell adhesion, which was already present after 24 h of incubation at the concentration of 10 and 20 μM (Figure 2C). After a 48-h treatment, a significant inhibitory effect was already present at a lower dose (5 μM). Conversely, in SH-SY5Y cells no inhibitory effect was observed for all the tested RGZ concentrations, neither after a 24-h nor after a 48-h exposure (Figure 2D).

Effects of the PPARγ antagonist BADGE on cell proliferation and viability

In order to determine whether the effects of RGZ on cell proliferation and viability were truly mediated by PPARγ, additional experiments were performed in SK-N-AS using the PPARγ antagonist BADGE. The exposure to 20 μM BADGE did not determine any effect both on [³H]-thymidine incorporation and on cell viability, thus indicating that at this concentration this compound had no toxic effect. The same concentration of BADGE was able to revert the inhibitory effect of 20 μM RGZ on [³H]-thymidine incorporation (Figure 3A) as well as on cell viability (Figure 3B).

Effect of RGZ on cell invasiveness and on the expression of MMP-9/TIMP-1

The effect of RGZ (20 μM) on the invasiveness ability of NB cells was determined. When SK-N-AS cells were tested, the number of invasive cells markedly decreased after RGZ exposure. A representative example is shown in Figure 4A and B. At variance with SK-N-AS, no effect was observed in SH-SY5Y (not shown).

Effects of RGZ on apoptosis

In order to determine whether RGZ is also able to induce apoptosis in the two NB cell lines used in this study, the number of cells showing positivity for cleaved caspase 3 was evaluated by immunocytochemistry. In SK-N-AS, the amount of apoptotic cells in untreated samples was 0.44 ± 0.23% (mean ± s.e.), whereas RGZ treatment (20 μM) significantly increased the number of caspase-positive cells.
3-positive cells (2.12 ± 0.24%, P < 0.05). In SH-SY5Y, there was no difference in the amount of cleaved caspase 3-positive cells in untreated vs RGZ-treated samples (0.43 ± 0.13 vs 0.49 ± 0.11%, respectively, P > 0.05). Representative examples are shown in Figure 6. Several immunostained SK-N-AS cells were present after exposure to RGZ (Figure 6B), compared to the absence of staining in untreated cells (Figure 6A). In contrast, only a very few SH-SY5Y cells showed positivity for cleaved caspase 3 either before (Figure 6C) or after (Figure 6D) RGZ treatment. The effect of treatment with RGZ (20 μM) on caspase 3 activation in SK-N-AS was confirmed by flow cytometry analysis. After labelling activated caspase 3, treated cells showed a shift to higher values of fluorescence, when compared to control cells (9.73, treated vs 8.28, control, median values of fluorescence histograms). Conversely, no effect was observed in SH-SY5Y after RGZ exposure. Transfection of NB cell lines with a human PPARγ expression plasmid stimulated the reporter activity about 1.5- and eight-fold over the control in SK-N-AS and SH-SY5Y, respectively. Furthermore, the treatment of PPARγ-transfected cells with RGZ caused an additional five- and 17-fold increase of the reporter activity in the two cell lines, respectively.

PPARγ activity in NB cells

Because of the different response to RGZ, PPARγ activity was determined in SK-N-AS and SH-SY5Y cells. Peroxisome proliferator-activated receptor γ activity was investigated by transient transfection assay and PPARγ transactivation was monitored by the activity of transfected adipocyte response element-7-galactosidase reporter cells. In SK-N-AS, RGZ treatment (20 μM) induced a near three-fold increase of the reporter activity compared to untreated control cells (Figure 7). Conversely, no effect was observed in SH-SY5Y after RGZ exposure. Transfection of NB cell lines with a human PPARγ expression plasmid stimulated the reporter activity about 1.5- and eight-fold over the control in SK-N-AS and SH-SY5Y, respectively. Furthermore, the treatment of PPARγ-transfected cells with RGZ caused an additional five- and 17-fold increase of the reporter activity in the two cell lines, respectively.

PPARγ gene sequencing and expression in NB cells

In order to determine whether the different function of PPARγ as a transcriptional activator in the two NB cell lines was due to gene mutations, sequence analysis of the entire coding region of the PPARγ gene was performed. However, no mutation was found in either SH-SY5Y or SK-N-AS. Similarly, no evident difference in the amount of expression of PPARγ protein was observed between the two cell lines, as assessed by Western blot analysis (Figure 8A). Equivalent protein loading was verified by staining parallel gels with Coomassie R, as shown in Figure 8B. Because there is evidence that PPARγ phosphorylation reduces the activity of the receptor (Shao and Lazar, 1999), PPARγ immunoprecipitation was performed, followed by SDS–PAGE and Western blot analysis using either an anti-PPARγ or an anti-phosphoserine monoclonal antibody. Interestingly, the amount of phosphorylated PPARγ was found to be markedly lower in SK-N-AS, thus suggesting that the higher PPARγ activity in this cell line compared to SH-SY5Y may be due to a different phosphorylation status of the receptor (Figure 8C).
DISCUSSION

In the present study, the effect of the PPARγ ligand RGZ on two NB cell lines, which have a stromal (S) (SK-N-AS) and a neuroblast (N) (SH-SY5Y) phenotype (Ciccarone et al., 1989; Ross et al., 2003; Servidei et al., 2004), was addressed. We first demonstrated that cell proliferation was readily inhibited by RGZ in the S-type cell line, whereas this effect was less evident in the N-type cells. Cell viability was significantly reduced by RGZ in both cell lines, although a much stronger maximal inhibition was observed in SK-N-AS. The effects of RGZ on cell proliferation and viability were readily counteracted by the PPARγ antagonist BADGE, thus indicating that PPARγ activation has an important role in eliciting these biological effects. Furthermore, the capability of cultured cells to adhere to a solid substrate in response to RGZ was assessed. In fact, cell adhesion assay is an indicator of cell spreading and adhesion molecules play a critical role in promoting the processes leading to tumour invasion and metastasis (Cavallaro and Christofori, 2004). Again, we found that RGZ readily decreased the adhesion of SK-N-AS cells starting from a 24-h exposure (10 and 20 μM), whereas it had no effect on SH-SY5Y cells, even after a 48-h treatment. As an additional feature of malignant tumour cells, invasion is influenced by the expression/activity of MMPs in a PPARγ-dependent manner (Chantrain et al., 2004). MMPs have been addressed as a target for cancer therapy (Overall and Kleinfeld, 2004). RGZ has been shown for instance to reduce the expression/activity of MMPs in a PPARγ-independent manner in adrenal (Ferruzzi et al., 2005) and pancreatic tumour cells (Farrow et al., 2003; Galli et al., 2004) and to increase the expression of TIMP-1 in breast cancer cells (Liu et al., 2003). In SK-N-AS, but not in SH-SY5Y, RGZ significantly reduced the amount of MMP-9, whereas the expression of the MMP inhibitor TIMP-1 showed a small but not statistically significant increase at higher levels. The demonstration that the proliferation of embryonic stem cells with a null mutation for PPARγ could be inhibited by glitazones (Palakurthi et al., 2001), which were also able to differentiate these cells into macrophages (Chawla et al., 2001), further supports the presence of PPARγ-independent mechanisms of agonist action. Furthermore, in our study the role of RGZ in inducing apoptosis in NB cells was addressed. Interestingly, very recently apoptosis has been indicated as a surveillance mechanism against metastases in NB (Stupack et al., 2006). In fact, it has been shown that suppression of caspase 8 expression occurs during the development of NB metastases in vivo, whereas reconstitution of caspase 8 expression potentiated apoptosis and counteracted the spreading of tumour cells. Peroxisome proliferator-activated receptor γ agonists induce apoptosis in different cell types and in most studies it has been shown that caspase 3 plays an important role in mediating apoptotic death (Grommes et al., 2004). We found that RGZ (20 μM) increased the amount of cleaved (i.e. activated) caspase 3 only in SK-N-AS cells. However, both immunocytochemistry and flow cytometry experiments did not show a very strong apoptotic effect of RGZ, thus suggesting that RGZ does not appear to play a major role in stimulating apoptosis in these NB cells. These results are partially in agreement with a previous report, in which RGZ (up to 50 μM) was not able to induce apoptosis in different NB cell lines, although SK-N-AS cells were not evaluated (Servidei et al., 2004).

The different response of SK-N-AS and SH-SY5Y cells to RGZ appeared to be due to the function of PPARγ as a transcriptional activator, which was readily demonstrated in SK-N-AS but not in SH-SY5Y, as assessed by transient transfection with a PPARγ-responsive element. This difference might be due to a mutation of the PPARγ gene. No mutation in exons 3 and 5, the biologically important regions for DNA and lipid binding of PPARγ, respectively, has been detected previously in the two NB cell lines,

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**Figure 5** Amount of MMP-9 mRNA (A) and protein (B) and of TIMP-1 mRNA (C) in SK-N-AS cells in the absence or in the presence of different concentrations of RGZ. *P<0.05 vs control (indicated as C).
which were used in our study (Servidei et al, 2004). A K422Q mutation has been described previously in exon 6 of PPARγ in colon cancer cell lines, but it did not alter ligand-induced transactivation (Gupta et al, 2003). Nevertheless, we sequenced the entire coding region of the PPARγ gene in SK-N-AS and in SH-

![Image of SK-N-AS and SH-SY5Y cells](image)

**Figure 6** Immunocytochemistry for the detection of cleaved caspase 3-positive cells. (A and C) Control untreated cells (SK-N-AS and SH-SY5Y, respectively). (B and D) RGZ-treated cells (SK-N-AS and SH-SY5Y, respectively). Magnification × 40.

**Figure 7** PPARγ transcriptional activity in control untreated NB cells (C), in cells treated with RGZ (20 μM) and in cells transfected with PPARγ in the absence or in the presence of RGZ. L/C: peroxisome proliferator response element-n73-tk-luciferase reporter activity, normalised for CAT activity. *P<0.05 vs C. **P<0.05 vs PPARγ transfected cells in the absence of RGZ.

![Graph showing transcriptional activity](image)

**Figure 8** (A) Western blot analysis of the amount of PPARγ in SK-N-AS and SH-SY5Y. PANC-1 and HPAC cells (human pancreatic adenocarcinomas) were used as the positive and negative control, respectively. (B) Coomassie R-stained gel, showing equal protein loading of the same samples indicated in A. (C) Detection of total (anti-PPARγ antibody) and phosphorylated (anti-P-Ser antibody) PPARγ, after PPARγ immunoprecipitation.

![Western blot images](image)

SYSY and no mutation was found. Only when SH-SY5Y cells were transfected with a human PPARγ expression plasmid, the response to RGZ was effectively elicited, thus confirming that the lack of
response was truly due to the very low transactivation potential of the endogenous PPARγ. It has to be mentioned that the phosphorylated status of PPARγ is important in determining its transcriptional activity. In fact, there is evidence that PPARγ phosphorylation reduces the activity of the receptor (Shao and Lazar, 1999). We performed Western blot analysis of total and phosphorylated PPARγ and we found that the amount of the phosphorylated form was markedly lower in SK-N-AS than in SH-SY5Y. This finding indicates that the lower activity of PPARγ in the latter NB cell line may be due to a different phosphorylation status of the receptor. In addition, it might be hypothesised that PPARγ overexpression in transfected SH-SY5Y cells might increase the level of unphosphorylated PPARγ beyond a critical threshold, thus eliciting transcriptional activity. Other recent studies have shown variable antineoplastic effects of PPARγ agonists in NB cell lines expressing PPARγ (Han et al, 2001, Servidei et al, 2004, Valentinier et al, 2005), but the activity of PPARγ was addressed only in one study, in which PPARγ was found to be functionally active in different cell lines and particularly in SK-N-AS (Servidei et al, 2004). However, the relationship between PPARγ transactivation and the response to TZDs was not addressed. Thus, our study extends the work of other authors, because the different response of NB cells to TZDs was related for the first time to differences in PPARγ transactivation, which, in turn, appear to be due to a different phosphorylation status of the receptor.

REFERENCES
Benvenuti S, Luciani P, Vannelli GB, Gelmini S, Franceschi E, Serio M, Peri A (2005) Estrogen and selective estrogen receptor modulators exert neuroprotective effects and stimulate the expression of selective Alzheimer’s disease indicator-1, a recently discovered antiapoptotic gene, in human neuroblast long-term cell cultures. J Clin Endocrinol Metab 90: 1775 – 1782
Burstein HJ, Demetri GD, Mueller E, Sarraf P, Spiegelman BM, Winer EP (2003) Use of the peroxisome proliferator-activated receptor (PPAR) gamma ligand troglitazone as treatment for refractory breast cancer: a phase II study. Breast Cancer Res Treat 79: 391 – 397
Carloni V, Romanelli R, Morrici AM, Pinzani M, Laffi G, Crotogini G, Gentilini P (1998) Knockout of alphabeta-talin-integrin expression reverses the transformed phenotype of hepatocarcinoma cells. Gastroenterology 115: 433 – 442
Cavallaro U, Christofori G (2004) Multitasking in tumor progression: signaling functions of cell adhesion molecules. Ann NY Acad Sci 1014: 58 – 66
Chantrain CF, Shimada H, Jodele S, Groshen S, Yeku A, Shalinsky DR, Werb Z, Coussens LM, DeClerck YA (2004) Stromal matrix metalloproteinase-9 regulates the vascular architecture in neuroblastoma by promoting pericyte recruitment. Cancer Res 64: 1675 – 1686
Chawla A, Barak Y, Nagy L, Diao D, Tontonoz P, Evans RM (2001) PPAR-gamma dependent and independent effects on macrophage gene expression in lipid metabolism and inflammation. Nat Med 7: 48 – 52
Ciccaveco V, Spengler BA, Meyers MB, Biedler JL, Ross RA (1989) Phenotypic diversification in human neuroblastoma cells: expression of distinct neural crest lineages. Cancer Res 49: 219 – 225
Cioppa F, Simi L, Luciani P, Petraglia F, Susini T, Cobelli L, Serio M, Maggi M, Peri A (2004) Expression of uteroglobin and matrix metalloproteinase-9 genes in endometrial cancer: relationship to estrogen and progesterone receptor status. Oncol Rep 11: 427 – 433
Costa-Guda J, Rosen ED, Jensen RT, Chung DC, Arnold A (2005) Mutational analysis of PPARγ as a candidate tumour suppressor gene in enteropancreatic endocrine tumours. Clin Endocrinol 62: 603 – 606
Cox PJ, Ryan DA, Hollis FJ, Harris AM, Miller AK, Vosden M, Cowley H (2000) Absorption, disposition, and metabolism of rosiglitazone, a potent thiazolidinedione insulin sensitizer, in humans. Drug Metab Dispos 28: 772 – 780
Crabb DW, Dixon JE (1987) A method for increasing the sensitivity of chloramphenicol acetyltransferase assay in extracts of transfected cultured cells. Anal Biochem 63: 88 – 92
de Wet JR, Wood KV, DeLuca M, Helinski DR, Subramani S (1987) Firefly luciferase gene: structure and expression in mammalian cells. Mol Cell Biol 7: 725 – 737
Debrock G, Vanhentenrijk V, Sciot R, Debiec-Rychter M, Oyen R, Van Oosterom A (2003) A phase II trial with rosiglitazone in liposarcoma patients. Br J Cancer 89: 1409 – 1412
Demetri GD, Fletcher CD, Mueller E, Sarraf P, Naujoks R, Campbell N, Spiegelman BM, Singer S (1999) Induction of solid tumor differentiation by the peroxisome proliferator-activated receptor-γ ligand troglitazone in patients with liposarcoma. Proc Natl Acad Sci USA 96: 3951 – 3956
Desvergne B, Wahli W (1999) Peroxisome proliferator-activated receptors: nuclear control of metabolism. Endocr Rev 20: 649 – 658
Farrow B, O’Connor KL, Hashimoto K, Iwamura T, Evers BM (2003) Selective activation of PPARγ-gamma inhibits pancreatic cancer invasion and decreases expression of tissue plasminogen activator. Surgery 134: 206 – 212
Ferruzzi P, Ceni E, Tarocchi M, Grappone C, Milani S, Galli A, Fiorelli G, Serio M, Mannelli M (2005) Thiazolidinediones inhibit growth and invasiveness of the human adrenocortical cancer cell line H295R. J Clin Endocrinol Metab 90: 1332 – 1339
Galli A, Ceni E, Crab DW, Mello T, Salzano R, Grappone C, Milani S, Surrenti E, Surrenti C, Casini A (2004) Antidiabetic thiazolidinediones inhibit invasiveness of pancreatic cancer cells via PPARγ independent mechanisms. Gut 53: 1688 – 1697
Gamble JR, Vadas MA (1988) A new assay for the measurement of the attachment of neutrophils and other cell types to endothelial cells. J Immunol Methods 109: 175 – 184
Garaventa A, Lukrach R, Lo Piccolo MS, Cavadini E, Montaldo PG, Pizziola MR, Boni L, Ponzoni M, Decensi A, De Bernardi B, Bellani FF, Formelli F (2003) Phase I trial and pharmacokinetics of fenretinide in children with neuroblastoma. Clin Cancer Res 9: 2032 – 2039
Grommes C, Landreth GE, Heneka MT (2004) Antineoplastic effects of peroxisome proliferator-activated receptor γ agonists. Lancet Oncol 5: 419 – 429
Gupta RA, Sarraf P, Mueller E, Brockman JA, Prusakiewicz JJ, Eng C, Willson TM, DuBois RN (2003) Peroxisome proliferator-activated receptor-γ-mediated differentiation. J Biol Chem 278: 22669 – 22677
Han SW, Greene ME, Pitts J, Wada RK, Sidell N (2001) Novel expression and function of peroxisome proliferator-activated receptor gamma (PPARγ) in human neuroblastoma cells. Clin Cancer Res 7: 98 – 104
Hisatake J, Ikezoe T, Carey M, Holden S, Tomoyasu S, Koehler HP (2000) Down-regulation of prostate-specific antigen expression by ligands for...
Reginato MJ, Bailey ST, Krakow SL, Minami C, Ishii S, Tanaka H, Lazar MA (1998) A potent antidiabetic thiazolidinedione with unique peroxisome proliferator-activated receptor γ-activating properties. J Biol Chem 273: 32679 – 32684

Reynolds CP, Matthay KK, Villalbanca JG, Maurer BJ (2003) Retinoid therapy of high-risk neuroblastoma. Cancer Lett 197: 185 – 192

Ross RA, Biedler JL, Spangler BA (2003) A role for distinct cell types in determining malignancy in human neuroblastoma cell lines and tumors. Cancer Lett 197: 35 – 39

Sarraf P, Mueller E, Smith WM, Wright HM, Kum JB, Aaltoenen LA, de la Chapelle A, Spiegelman BM, Eng C (1999) Loss-of-function mutations in PPAR γ associated with human colon cancer. Mol Cell 3: 799 – 804

Servidei T, Morosetti R, Ferlini C, Cusano G, Scambia G, Mastrangelo R, Koeffler HP (2004) The cellular response to PPARγ ligands is related to the phenotype of neuroblastoma cell lines. Oncol Res 14: 345 – 354

Shao D, Lazar MA (1999) Modulating nuclear receptor function: may the phos be with you. J Clin Invest 12: 1617 – 1618

Shimada H, Joshi VV (1997) Pathology of neuroblastoma. In Principles and Practice of Genitourinary Oncology Raghavan D, Scher HI, Leibel SA, Lange H (eds). pp 1069 – 1076. New York: Lippincott-Raven Publishers

Strakova N, Ehrmann J, Dzubak P, Bouchal J, Kolar Z (2004) The synthetic ligand of peroxisome proliferator-activated receptor-γ ciglitazone affects human glioblastoma cell lines. J Pharmacol Exp Ther 309: 1239 – 1247

Stupack DG, Teitz T, Potter MD, Mikolol D, Houghton PJ, Kidd VJ, Lahti JM, Cheres DA (2006) Potentiation of neuroblastoma metastasis by loss of caspase-8. Nature 439: 95 – 99

Sugiura Y, Shimada H, Seeger RC, Laug WE, DeClerck YA (1998) Matrix metalloproteinases-2 and -9 are expressed in human neuroblastoma: contribution of stromal cells to their production and correlation with metastasis. Cancer Res 58: 2209 – 2216

Valentinier U, Carlsson M, Ertemann R, Hildebrandt H, Schumacher U (2005) Ligands for the peroxisome proliferator-activated receptor-γ have inhibitory effects on growth of human neuroblastoma cells in vitro. Toxicology 213: 157 – 168

Voigt A, Zintl F (2003) Effects of retinoic acid on proliferation, apoptosis, citotoxicity migration, and invasion of neuroblastoma cells. Med Pediatr Oncol 40: 205 – 213

Yki-Järvinen H (2004) Thiazolidinediones. N Engl J Med 351: 11016 – 11018