PNU-91325 increases fatty acid synthesis from glucose and mitochondrial long chain fatty acid degradation: a comparative tracer-based metabolomics study with rosiglitazone and pioglitazone in HepG2 cells

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The mitochondrial membrane protein termed “mitoNEET,” is a putative secondary target for insulin-sensitizing thiazolidinedione (TZD) compounds but its role in regulating metabolic flux is not known. PNU-91325 is a thiazolidinedione derivative which exhibits high binding affinity to mitoNEET and lowers cholesterol, fatty acid and blood glucose levels in animal models. In this study we report the stable isotope-based dynamic metabolic profiles (SIDMAP) of rosiglitazone, pioglitazone and PNU-91325 in a dose-matching, dose-escalating study. One and 10 \( \mu \text{M} \) concentrations 1 and 10 \( \mu \text{M} \) drug concentrations were introduced into HepG2 cells in the presence of either \([1,2\text{-}^{13}\text{C}]\text{-glucose}\) or \([U\text{-}^{13}\text{C}]\text{stearate}\), GC/MS used to determine positional tracer incorporation (mass isotopomer analysis) into multiple metabolites produced by the Krebs and pentose cycles, \textit{de novo} fatty acid synthesis, long chain fatty acid oxidation, chain shortening and elongation. Rosiglitazone and pioglitazone (10 \( \mu \text{M} \)) increased pentose synthesis from \([U\text{-}^{13}\text{C}]\text{stearate}\) by 127% and 185%, respectively, while PNU-91325 rather increased glutamate synthesis in the Krebs cycle by 113% as compared to control vehicle treated cells. PNU-91325 also increased stearate chain shortening into palmitate by 59%. Glucose tracer-derived \textit{de novo} palmitate and stearate synthesis were increased by 1 and 10 \( \mu \text{M} \) rosiglitazone by 41% and 83%, respectively, and by 63% and 75% by PNU-91325. Stearate uptake was also increased by 10 \( \mu \text{M} \) PNU-91325 by 15.8%. We conclude that the entry of acetyl Co-A derived from long-chain fatty acid \( \beta \)-oxidation into the mitochondria is facilitated by the mitoNEET ligand PNU-91325, which increases glucose-derived long chain fatty acid synthesis and breakdown via \( \beta \)-oxidation and anaplerosis in the mitochondria.

KEY WORDS: PNU-91325; mitoNEET; stable isotope-based dynamic metabolic profiling (SIDMAP); fatty acid oxidation; glucose metabolism; diabetes.

1. Introduction

Thiazolidinedione (TZD) compounds are used for the treatment of type 2 diabetes and are believed to exert their beneficial effects by increasing glucose-derived fatty acid synthesis secondary to the increased degradation of long chain fatty acids through the activation of the nuclear receptor peroxisome proliferator-activated receptor-gamma (PPAR\textgamma). Studies with a tritiated photo-affinity labeled pioglitazone (1) analog identified a novel saturable TZD binding site also in mitochondria (Colca \textit{et al.}, 2004). The binding correlated with a \(<17\text{-kDa}\text{protein}\), which showed selectivity of competition by analogs. The protein is termed “mitoNEET,” and has been found in the mitochondrial fraction of rodent brain, liver, and skeletal muscle. The protein also exists in low levels in preadipocytes, but expression increases exponentially in differentiated adipocytes (Colca \textit{et al.}, 2004).

Several pioglitazone (1) derivatives have been tested as antidiabetic TZDs for possible clinical development (Tanis \textit{et al.}, 1996). PNU-91325 (2)

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was found to be the most effective TZD and proved to have greater binding to mitoNEET. Thus modification of mitoNEET function may, at least in part, contribute to lowering circulating lipid levels and glucose in the KKA\textsuperscript{Y} mouse model of human type-2 diabetes. Evidence has recently been presented suggesting that the insulin sensitizing effects of thiazolidinediones may correlate with modification of mitochondrial function rather
than activation of PPARγ (Colca, 2006). Changes in mitochondrial function may alter the usage of cellular metabolites (e.g., sugars and fatty acids) in a way that favors insulin sensitivity.

In this study we report the stable isotope-based dynamic metabolic profile (SIDMAP) of rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) in a dose-matching, dose-escalating in vitro study. In order to follow substrate flow in the glucose metabolic network and the fate of the long chain fatty acid derived acyl-CoA we used two tracers; [1,2-13C2]-d-glucose or [U-13C18] stearate pool in differentiated primary hepatocellular carcinoma (HepG2) cells and gas chromatography/mass spectrometry (GC/MS) to analyze RNA ribose synthesis, lactate production, TCA cycle glutamate synthesis, as well as myristate, palmitate, stearate and oleate synthesis and turnover using the comprehensive multiple pathway tracer substrate flow analysis platform of the SIDMAParray (Boros et al., 2004). It is herein demonstrated that PNU-91325 (2) potently increases de novo long chain fatty acid synthesis from glucose, while it also increases mitochondrial breakdown of long-chain derived acyl-CoA via TCA cycle anaplerosis and glutamate synthesis. On the contrary, rosiglitazone (3) and pioglitazone (1) are more effective in re-routing stearate derived acyl-CoA towards glyconeogenesis and the synthesis of five carbon metabolites such as ribose in the pentose cycle. We conclude that the mitoNEET protein is a secondary TZD target and has a role in increasing mitochondrial long chain fatty acid derived acyl-CoA disposal through TCA cycle anaplerosis.

2. Materials and methods

Two non-radiating stable isotope tracer carbon labeled substrates were used in this metabolic flux analysis study: (a) [1,2-13C2]-d-glucose was purchased with >99% purity and 99% isotope enrichment for each position from Cambridge Isotope Laboratories, Inc. (Andover, MA, USA); and (b) [U-13C18] stearic acid (Spectra Stable Isotopes, Spectra Gases Inc., Branchburg, NJ). Rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) were provided by Pfizer, Inc., under a material transfer agreement.

2.1. Cells and cell culture

Human liver hepatocellular (epithelial) carcinoma HepG2 cells (ATCC CRL11997) were purchased from American Type Culture Collection (ATCC, Manassas, VA, USA). HepG2 cells have an average doubling time of 34 h in DMEM with 10% fetal bovine serum and 2.5% horse serum (Gibco/BRL, Gaitersburg, MD) in the presence of antibiotics. The cells were incubated at 37°C, 5% CO2 and 95% humidity and passed by using trypsin 0.25% (Gibco/BRL) no more than three times after receipt from the ATCC and prior to use in this study. HepG2 cells have previously been used in both in vitro and in vivo experiments and responded to treatment with troglitazone and other compounds with characteristic metabolic profile changes showing altered macromolecule synthesis and fatty acid cycling (Lee et al., 1998a). Tracer-labeled parallel cultures of HepG2 cells were used in this study as controls to compare vehicle (DMSO) treated cell cultures to rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) treatment in an escalating regime of drug treatment.

Seventy-five percent confluent cultures of HepG2 cells were incubated in either [1,2-13C2]-d-glucose or [U-13C18] stearate-containing media (100 mg/dl total glucose concentration = 5 mM; 50% isotope enrichment; or 0.5 mM uniformly labeled stearate). Cells were plated at a density of 106 per T75 culture flask and rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) were added in two concentrations, 1 or 10 μM dissolved in 10% DMSO – 90% culture media. Control cultures were treated with vehicle (10% DMSO) only. The doses of rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) for the present study were selected based on in vitro experiments demonstrating that these drugs effectively control glucose levels via PPARγ activation in various human cell culture systems in the 1 to a 10 μM dose range (for review: Otto et al., 2002). Drug treatments were carried out for 72 h in the presence of each tracer. Glucose and U-stearate 13C tracer levels in the medium were monitored using a Cobas Mira chemistry analyzer (Roche Diagnostics, Pleasanton, CA, USA). Fresh, tracer enriched culture medium and drug treatments were applied every 24 h, three times during the experiment, in order to provide for the cells with steady tracer-enriched medium and drug doses throughout the 72 h experimental period. After 72 h the cells were scraped, spun and frozen at −80°C until further processing as described below.

2.2. RNA ribose stable isotope studies

For mass isotopomer analysis RNA ribose was isolated by acid hydrolysis of cellular RNA after Trizol (Invitrogen Life Technologies, Carlsbad, CA, USA) extraction from the cell pellets. Ribose was derivatized to its aldonitrile acetate form using hydroxylamine in pyridine with acetic anhydride (Supelco, Bellefonte, PA, USA) before mass spectral analyses. We monitored the ion cluster around the m/z 2256 (carbons 1–5 of ribose) (chemical ionization, CI) and m/z 2217 (carbons 3–5 of ribose) and m/z 242 (carbons 1–4 of ribose) (electron
impact ionization, EI) to determine molar enrichment and the positional distribution of $^{13}$C in ribose. By convention, the base mass of $^{13}$C compounds (with their derivatization agents) is given as $m_0$ as measured by mass spectrometry as described elsewhere (Katz et al., 1993). Ribose molecules labeled with a single $^{13}$C atom on the first carbon position ($m_1$) recovered from RNA were used to gauge the ribose fraction produced by direct oxidation of glucose through the G6PD pathway. Ribose molecules labeled with $^{13}$C on the first two carbon positions ($m_2$) were used to measure the fraction produced by transketolase. Doubly labeled ribose molecules ($m_2$ and $m_4$) on the fourth and fifth carbon positions were used to measure molar fraction produced by triose phosphate isomerase and transketolase (Boros et al., 2002). Because transketolase has the highest metabolic control coefficient in the non-oxidative branch of the pentose cycle (Sabate et al., 1995), we use the term transketolase throughout the paper. It should be noted, though, that transketolase and transaldolase, besides other enzymes, all are participants in non-oxidative pentose cycle metabolism in human cells.

2.3. Lactate

Lactate from the cell culture media (0.2 ml) was extracted by ethylene chloride after acidification with HCl. Lactate was derivatized to its propylamine-heptafluorobutyrate ester form and the $m/z$328 (carbons 1–3 of lactate) (chemical ionization, CI) was monitored for the detection of $m_1$ (recycled lactate through the PC) and $m_2$ (lactate produced by the Embden–Meyerhof–Parnas pathway) for the estimation of pentose cycle activity (Lee et al., 1998b). In this study we recorded the $m_1/m_2$ ratios in lactate produced and released by HepG2 cells in order to determine pentose cycle activity and pentose cycling versus anaerobic glycolysis in response to 1 and 10 μM rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) treatments.

2.4. Glutamate

$^{13}$C label distribution in glutamate (glutamic acid), which is synthesized from α-keto-glutaric acid of the TCA cycle, from tracer labeled glucose or stearate is suitable for determining mitochondrial glucose and fatty acid disposal via anabolic substrate use within the cycle, also known as anaplerotic flux. For glutamate extraction tissue culture medium (1 ml) was first treated with 1 ml of 6% perchloric acid and the supernatant was passed through a 3 cm³ Dowex-50 (H⁺) column. Amino acids were eluted with 15 ml 2 N ammonium hydroxide. In order to separate glutamate from glutamine, the amino acid mixture was passed through a 3 cm³ Dowex-1 (acetate) column, and then collected with 15 ml 0.5 N acetic acid. The glutamate fraction from the culture medium was converted to its trifluoroacetyl butyl ester (TAB) (Leimer et al., 1977). Under EI conditions, ionization of TAB-glutamate produces two fragments, $m/z$198 and $m/z$152, corresponding to C2–C5 and C2–C4 of glutamate. Glutamate labeled on the 4–5 carbon positions indicates pyruvate dehydrogenase activity while glutamate labeled on the 2–3 carbon positions indicates pyruvate carboxylase activity for the entry of glucose carbons to the TCA cycle. TCA cycle anabolic glucose utilization is calculated based on the $m_1/m_2$ ratios of glutamate (Lee et al., 1996).

2.5. Lipid extraction and analysis

Lipid extractions were performed using methods described elsewhere (Lowenstein et al., 1975). In brief, the Trizol extract (bottom layer) of cell pellets after RNA extraction were saponified with 500 μl of 30% KOH–ethanol (1:1 v/v) at 70 °C overnight. The aqueous phase was acidified, and fatty acids were extracted in petroleum ether and dried under a stream of nitrogen. Fatty acids were methylated with 0.5 ml 0.5 N HCl in methanol for GC/MS analysis. Palmitate, stearate and oleate were monitored at $m/z$270, $m/z$298 and $m/z$264, respectively, with the enrichment of $^{13}$C labeled acetyl units either from tracer labeled glucose or stearate, which reflect synthesis, elongation and desaturation of the new lipid fraction as determined by mass isotopomer distribution analysis (MIDA) of different isotopomers (Lee et al., 1991). Mass isotopomer distribution was determined using the method of Lee et al., which corrects for the contribution of derivatizing agent and $^{13}$C natural abundance to the mass isotopomer distribution of the compound of interest. The calculated mass isotopomer distribution is expressed as molar fractions ($m_0$, $m_1$, $m_2$, $m_3$, etc.), which are the fractions of molecules containing 0, 1, 2, 3,... $^{13}$C substitutions, respectively. Data reduction and regression analyses were performed using the computer software Microsoft Excel™ version 5.0.

2.5.1. Determination of precursor enrichment

β-oxidation of [U-$^{13}$C₁₈]stearate generates [1,2-$^{13}$C₂] acetyl-CoA, which is subsequently used as a precursor for de novo lipogenesis, chain elongation, energy production and anaplerosis via the tricarboxylic acid cycle. Thus, β-oxidation can be assessed through the determination of [1,2-$^{13}$C₂]acetyl-CoA enrichment in the different synthetic products.

2.5.2. Acetyl-CoA enrichment for de novo synthesis from glucose

The enrichment of the cytosolic acetyl-CoA pool used in de novo fatty acid synthesis was determined from the mass isotopomer distribution in palmitate and stearate using the [1,2-$^{13}$C₂]-β-glucose as the tracer. De novo synthesis produces these long chain saturated fatty acids with 2, 4, or 6 $^{13}$C atoms (m2, m4, and m6). The
distribution of these mass isotopomers has been previously shown to be a binomial distribution (Hellerstein, 1991; Lee et al., 1992). Thus, the acetyl-CoA enrichment may be obtained from the consecutive mass isotopomer ratio m4/m2 = (n–1)p/q = 3.5p/q, where n is the number of acetyl units in palmitate = 8, p is the enrichment of [1,2–13C2]-acetyl-CoA, q is the unenriched acetyl-CoA, and p + q = 1.

2.5.3. Acetyl-CoA enrichment from mitochondrial oxidation

[1,2–13C2]-acetyl-CoA produced either from [1,2–13C2]-d-glucose or [U–13C18]stearate from β-oxidation combines with oxaloacetic acid and forms citric acid in the mitochondria. Citrate participates in the tricarboxylic acid cycle, eventually forming [α-4,5–13C2]ketoglutarate and glutamate (Haber and Lapidot, 2001). The enrichment of the m2 component of the C2–C5 fragment generally reflects tricarboxylic acid cycle activity. However, it is well known that contribution from unlabeled glutamate from the cells in culture with glutamine-containing medium can substantially dilute the glutamate enrichment (Darmaun et al., 1988). Therefore, the enrichment of the [4,5–13C2]glutamate reflects the lower limit of mitochondrial acetyl-CoA cycling and anaplerosis.

2.5.4. Acetyl-CoA enrichment for chain elongation

Enzymes involved in chain elongation from [U–13C18]stearate to longer chain fatty acids are located in the endoplasmic reticulum, mitochondria, and peroxisomes (Vance and Vance, 2002). Chain elongation of uniformly labeled stearate with [1,2–13C2]acetyl-CoA produces characteristic m + 18 clusters that can be used to determine the acetyl-CoA enrichment according to the rules of combination of two labeled precursors. The ratio of m + 2 to m + 0 acetyl-CoA for chain elongation is given by the ratio of m + 20 to m + 18 in arachidate using the [U–13C18]stearate tracer. The enrichment of [1,2–13C2]acetyl-CoA for each step of chain elongation was determined using the method described elsewhere using HepG2 cells as the in vitro model of fatty acid metabolism in the presence of [U–13C18]stearate (Wong et al., 2004).

2.6. Gas chromatography/mass spectrometry (GC/MS)

Mass spectral data were obtained on the HP5973 mass selective detector connected to an HP6890 gas chromatograph. The settings were as follows: GC inlet 250 °C, transfer line 280 °C, MS source 230 °C, MS Quad 150 °C. An HP-5 capillary column (30 m length, 250 μm diameter, 0.25 μm film thickness) was used for glucose, ribose, deoxyribose, glutamate and lactate analyses. Alternatively, a Bpx70 column (25 m length, 220 μm diameter, 0.25 μm film thickness, SGE Incorporated, Austin, TX) was used for fatty acid analysis with specific temperature programming for myristate (C:14), palmitate (C:16), stearate (C:18), oleate (C:18–1), as well as C:20, C:22 and C:24 fatty acid analyses.

2.7. Data analysis and statistical methods

Each experiment was carried out using triplicate cell cultures for each condition within each experiment and experiments were repeated once. Mass spectroscopic analyses were carried out by three independent automatic injections of 1 μl samples by the automatic sampler and accepted only if the standard sample deviation was less than 1% of the normalized peak intensity. Statistical analysis was performed using the Student’s t-test for unpaired samples. Two-tailed significance at the 99% confidence interval (μ ± 2.58 σ, p < 0.01) indicated significant differences in glucose carbon metabolism in control and rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) treated HepG2 cells. SIDMAParray™ was produced by MeTa Informatics (San Diego, CA).

3. Results

In this study the stable isotope-based metabolic profiles of rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) treated HepG2 cells are presented using two tracers, [1,2–13C2]-d-glucose and [U–13C18]stearic acid in separate experiments. The metabolic profiles reported in the SIDMAPArrays present tracer carbon flow changes in all major metabolic pathways, which include glycolysis, the pentose and TCA cycles, de novo fatty acid synthesis, chain elongation, chain shortening and desaturation, in a color scheme similar to gene expression studies.

The SIDMAParray using the [1,2–13C2]-d-glucose tracer is shown in figure 1. It can be seen from the array represented in figure 1 that rosiglitazone (3) and PNU-91325 (2) induced a significant increase in de novo fatty acid synthesis from glucose, including myristate, palmitate and stearate (box 3), while substrate flux through the pentose cycle producing ribose and deoxyribose were less affected by either drug (boxes 1 & 2). Rosiglitazone (3) and PNU-91325 (2) increased de novo stearate synthesis from glucose by 41% and 83% and by 63% and 75% using 1 and 10 μM doses, respectively. It can also be seen from the array that all compounds increased glucose-derived acetyl-CoA 13C enrichment of myristate and stearate. There were no significant changes in pentose cycle direct glucose oxidation and recycling into lactate or TCA cycle anaplerotic flux for the synthesis of glutamate from glucose (data not shown). It can be concluded from the array that 1 and 10 μM rosiglitazone (3) and PNU-91325 (2) induced similar changes in glucose carbon utilization in HepG2 cells, which was consistent with increased de novo long chain fatty acid synthesis from glucose, with little or no effect on the rate of glucose oxidation (13CO2 release in the TCA and
De novo release [Pentose and TCA cycles (\(\Delta^{13}C/^{12}C\))]

13C labeled fraction (\(\Sigma m\))

Total 13C enrichment (\(\Sigma_m\))

Synthesis via G6PDH (m1/\(\Sigma m\))

Myristate (C:14) Acetyl-CoA enrichment

Palmitate (C:16) Acetyl-CoA enrichment

Stearate (C:18) Acetyl-CoA enrichment

Myristate (C:14) Fraction of New Synthesis

Palmitate (C:16) Fraction of New Synthesis

Stearate (C:18) Fraction of New Synthesis

RNA ribose

DNA deoxyribose

Fatty acids

Figure 1. Stable isotope-based dynamic metabolic profile array (SIDMAParray) of rosiglitazone, pioglitazone and PNU-91325 in HepG2 cells using [1,2-\(^{13}\)C]-\(\alpha\)-glucose as the tracer. Treatments are indicated as follows: C = Control; 1R = 1 \(\mu\)M rosiglitazone; 10R = 10 \(\mu\)M rosiglitazone; 1P = 1 \(\mu\)M pioglitazone; 10P = 10 \(\mu\)M pioglitazone; 1PN = 1 \(\mu\)M PNU-91325; and 10PN = 10 \(\mu\)M PNU-91325. The color of each square corresponds to \(^{13}\)C tracer glucose flux via various synthetic pathways of ribose, deoxyribose and fatty acids as percent of control-DMSO treated cultures. A color scheme indicating the intensity of changes in substrate flux is provided in the bottom of the figure. Significant increase in de novo fatty acid synthesis from glucose are underlined and italicized (G6PDH = glucose 6-phosphate dehydrogenase; TPI = Triose-phosphate isomerase).

1. Experimental Procedure

2. The main difference in stearate utilization among the drugs is that 1 and 10 \(\mu\)M rosiglitazone (3) and pioglitazone (1) induced a dose dependent increase in pentose cycle 13C enrichment by 101\%, 127\% and 250\%, 285\% (box 1), while 1 and 10 \(\mu\)M PNU-91325 rather increased mitochondrial TCA cycle glutamate synthesis from stearate by 25\% and 113\% (box 2), respectively. Glutamate 13C enrichment from stearate was only increased using 10 \(\mu\)M rosiglitazone (3) or pioglitazone (1) by 22\% and 21\%, respectively. It appeared that the primary route by which stearate 13C tracer carbons entered the TCA cycle is via citrate synthesis, as 10 \(\mu\)M pioglitazone (2) and PNU-91325 (1) induced a 12\% and 34\% increase in glutamate synthesis via citrate (box 3). De novo palmitate synthesis from stearate-derived [13C]acetyl-CoA was increased by 102\%, while stearate chain shortening into palmitate was also increased by 59\% using 10 \(\mu\)M PNU-91325 (box 4).

3. Discussion

4. Metabolic profiling represents a logical development in the application of genomics, transcriptomics and proteomics in drug discovery and development. This study highlighted the relevance and robustness of stable isotope tracer-based dynamic metabolic profiles when applied to drugs discovered through reverse...
pharmacology (Harrigan et al., 2005). This study makes use of continued development of new approaches to generate and interrogate metabolome data sets. Here we have evaluated three thiazolidinediones that exhibit overall similar insulin sensitizing pharmacology in vivo but vary substantially in their ability to activate the transcription factor PPARγ. This study determined glucose and stearate flow redistribution among major metabolic pathways involving glycolysis, the pentose and TCA cycles, de novo fatty acid synthesis and degradation in response these three thiazolidinediones in living HepG2 primary liver tumor cells. The compounds tested in this study include two drugs marketed for the treatment of diabetes and a third analog that has potent insulin sensitizing activity without activating PPARγ-regulated gene transcription (Colca, 2006). Gene transcription analysis from earlier studies suggests that changes the expression of mitochondrial proteins might be a key factor in the insulin sensitizing pharmacology. Our findings using glucose as a metabolic tracer are that rosiglitazone (3), pioglitazone (1) and PNU-91325 (2) increase de novo synthesis of various fatty acids from glucose, and that PNU-91325 (2) surpasses the effect of both marketed drugs. Using stearate as a metabolic tracer it is obvious that rosiglitazone (3) and pioglitazone (1) primarily induce fatty acid breakdown by increasing glyconeogenesis and RNA pentose synthesis, while PNU-91325 (2) forces stearate-derived acetyl units into the mitochondria via citrate synthesis in a dose dependent manner.

The main implication of our SIDMAP study is that predominantly PPARγ and predominantly mitochondrially acting (mitoNEET ligand) compounds direct long chain fatty acid catabolism into different routes. PNU-91325 (2) is very effective in promoting mitochondrial stearate β-oxidation, while concomitantly increasing stearic acid uptake, chain shortening into palmitate and chain elongation into arachidonate. On the other hand, rosiglitazone (3) and pioglitazone (1) primarily increase pentose sugar synthesis from stearate, which indicates peroxisomal fatty acid metabolism and the production of acyl-CoA, which may enter the mitochondria and produce glutamate, but only at the high 10.0 μM dose in this dose matching study with PNU-91325. The potency of PNU-91325 (2) in increasing mitochondrial beta-oxidation of stearate and its disposal via glutamate is about 10 times higher than that of rosiglitazone (3) and pioglitazone (1). In previous in vivo experiments high resolution 2-D gel electrophoresis of livers showed that rosiglitazone (3) primarily produced changes in the expression pattern of many proteins involved in peroxisomal fatty acid β-oxidation (Edvardsson et al., 1999). Although rosiglitazone (3) and PNU-91325 (2) tend to increase the expression of many mitochondrial proteins and produce changes that favor fatty acid oxidation, PNU-91325 (2) does not significantly increase PPARγ-driven transcription either in vitro or in vivo (Colca, 2006). The most likely explanation for the effects of these compounds on gene expression and on the metabolic fluxes observed in these studies is that the overall effect of a given analog represents a blend of direct mitochondrial actions (e.g., binding to mitoNEET) and effects on other sites of binding (e.g., PPARγ).

It has been demonstrated that both PPARγ ligands investigated in this study, pioglitazone (1) and rosiglitazone (3) effectively decreased liver cancer formation using a diethylnitrosamine-induced liver cancer in Wistar rats (Guo et al., 2004) and colon carcinogenesis induced by azoxymethane in mice (Osawa et al., 2003). In our substrate flow tracer study, while rosiglitazone (3) and pioglitazone (1) increased RNA pentose synthesis from stearate, there was a decrease in tracer accumulation into DNA deoxyribose during the 72 h treatment period using these TZDs. On the other hand, DNA deoxyribobose synthesis was strongly suppressed by PNU-91325 (2) treatment from stearate. The glucose tracer study indicated unaltered rate of RNA and DNA ribose synthesis, which confirms the chemo-preventive effect of TZD based on direct substrate flow measurements via all arms of the pentose cycle.

We acknowledge two potential limitations of this study. First, because we studied primary hepatoma cells, it is not known how the studied compounds affect normal liver cell function and other tissue cells in a complex host. It is important to note that HepG2 cells grow rapidly in culture and they have higher specialized requirements for energy and macromolecule synthesis than primary cells. Conditions to test primary liver cells are under development. Metabolic labeling studies with primary cells are complicated by the fact that freshly isolated hepatocytes survive for 24 h providing limited time to produce a comprehensive SIDMAParray using either glucose or fatty acid tracers. In any case, HepG2 cells have maintained biotransformation properties of drugs and they also possess and express metabolic enzymes in a comparable manner with primary cultures of hepatocytes (Wilkening et al., 2003), while providing a highly viable and stable cell line for continuous tracer based metabolomics studies.

Second, it is currently not known what primary acyl-CoA species are produced from the tracer stearate substrate in response to the PPARγ agonist rosiglitazone (3) and pioglitazone (1) treatments. Based on assessing heavy incorporation of tracer carbons of stearate into the five carbon metabolite pool of ribose with no measurable tracer accumulation in lactate or palmitate it is likely that short chain fatty acids (C:4–C:6) are heavily produced and shuttled into the cytosol of HepG2 cells from the peroxisomes. As the use of C18–C20 chain elongation and other mass isotopomer analysis is the first quantitative method to determine peroxisomal acetyl-CoA enrichment (and therefore peroxisomal β-oxidation) (Wong et al., 2004), we will be able to identify those acyl-CoA pools in subsequent SIDMAP
studies. Such studies should make use of the tracer method for both peroxisomal oxidation and very long chain fatty acid synthesis in HepG2 cell cultures together with animal experiments.

It is a particularly significant finding that rosiglitazone (3) and pioglitazone (1) treatments strongly increase pentose formation from tracer stearate, thereby increasing NADPH production and peroxisomal function. In the mammalian liver many β- and per-oxidation related events require NADPH as the reducing equivalent. For example, peroxisomal β-oxidation produces hydrogen peroxide. Although hydrogen peroxide is not itself a radical, it is considered a reactive oxygen species because it is easily converted into hydroxyl radical, which has to be scavenged by NADPH-dependent reductases thus preventing cell damage. NADPH-dependent peroxidases are also important in the microsomal function of liver cells, therefore, increased carbon flow by rosiglitazone (3) and pioglitazone (1) towards the pentose cycle are expected during increased peroxisomal and microsomal fatty acid degradation (figure 2). In this paper we show direct evidence via net tracer carbon flow measurements from stearate towards pentoses that such link exists and such PPARγ ligands concomitantly increase peroxisomal degradation and pentose cycle metabolism in the same cells. These differences in metabolic flux from those produced by PNU-91325 (2) suggest that there will be some differences in the pharmacology produced by compounds that lack inherent PPARγ-like activity.

In summary, we evaluated glucose and fatty acid metabolism based on two separate metabolic tracer substrate experiments using cultured human primary liver carcinoma cells. Whatever the genetic and proteomic mechanisms are in the complex array of genes and proteins altered byTZDs, PNU-91325 (2) unquestionably generates a dramatically different substrate flow profile from the long chain fatty stearate than do rosiglitazone (3) and pioglitazone (1). PNU-91325 (2) directs β-oxidation of stearate and chain shortening by one acetyl unit resulting palmitate, then this acetyl unit participates in chain elongation and TCA cycle glutamate synthesis. Rosiglitazone (3) and pioglitazone (1) rather produce short chain fatty acids and their shortening is strongly linked with pentose synthesis (figure 3).

Based on our current findings, PNU-91325 (2) is a more potent inducer of mitochondrial fatty acid β-oxidation and TCA cycle anaplerosis than are the existing compounds to treat hyperlipidemic conditions in type 2 diabetes with desirable anti-proliferative effects and limited DNA deoxyribose synthesis in liver cells.

In order to reveal organ and tissue specific metabolic changes involving liver, muscle, adipose and other

Figure 2. Stable isotope-based dynamic metabolic profile array (SIDMAParray) of rosiglitazone, pioglitazone and PNU-91325 in HepG2 cells using [U-13C10]stearic acid as the tracer. Treatments are indicated as follows: C=Control; 1R =1 μM rosiglitazone; 10R =10 μM rosiglitazone; 1P =1 μM pioglitazone; 10P =10 μM pioglitazone; 1PN =1 μM PNU-91325; and 10PN =10 μM PNU-91325. The color of each square corresponds to 13C carbon flux from stearate via various synthetic pathways of ribose, deoxyribose, lactate, glutamate of the TCA cycle and fatty acids as percent of control-DMSO treated cultures. A color scheme indicating the intensity of changes in substrate flux is provided in the bottom of the figure. (DOR = deoxyribose; G6PDH = glucose 6-phosphate dehydrogenase; TPI = Triose-phosphate isomerase).
various cells, a tracer study in vivo is warranted using glucose and stearate tracer enriched diets in a wild type and diabetic host in order to better understand the effect of a primarily mitochondrially acting TZD (via mito-NEET?) plays in the regulation of long chain fatty acid breakdown and metabolism as compared to the currently marketed TZDs that are more potent PPAR\(\gamma\) activators.

Finally, it is evident that many reported effects of TZDs are clearly independent of PPARs. A most recent example is the demonstration that TZD inhibition of iNOS expression does not require PPAR involvement (Crosby et al., 2005). We suggest that many reported effects of TZD actions as being primarily mediated through PPAR should be interpreted with caution and that further investigation of the many secondary pharmacological effects associated with TZDs could contribute to the development of new therapeutic approaches to diabetes.

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