



Article Alternatively Spliced Landscape of PPARγ mRNA in Podocytes Is Distinct from Adipose Tissue

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Abstract: Podocytes are highly differentiated epithelial cells, and their structural and functional integrity is compromised in a majority of glomerular and renal diseases, leading to proteinuria, chronic kidney disease, and kidney failure. Traditional agonists (e.g., pioglitazone) and selective modulators (e.g., GQ-16) of peroxisome-proliferator-activated-receptor-y (PPARy) reduce proteinuria in animal models of glomerular disease and protect podocytes from injury via PPAR γ activation. This indicates a pivotal role for PPAR γ in maintaining glomerular function through preservation of podocytes distinct from its well-understood role in driving insulin sensitivity and adipogenesis. While its transcriptional role in activating adipokines and adipogenic genes is well-established in adipose tissue, liver and muscle, understanding of podocyte PPARy signaling remains limited. We performed a comprehensive analysis of PPARy mRNA variants due to alternative splicing, in human podocytes and compared with adipose tissue. We found that podocytes express the ubiquitous PPAR γ Var 1 (encoding γ 1) and not Var2 (encoding γ 2), which is mostly restricted to adipose tissue and liver. Additionally, we detected expression at very low level of Var4, and barely detectable levels of other variants, Var3, Var11, VartORF4 and Var9, in podocytes. Furthermore, a distinct podocyte vs. adipocyte PPAR-promoter-response-element containing gene expression, enrichment and pathway signature was observed, suggesting differential regulation by podocyte specific PPARy1 variant, distinct from the adipocyte-specific $\gamma 2$ variant. In summary, podocytes and glomeruli express several PPAR γ variants, including Var1 (γ 1) and excluding adipocyte-specific Var2 (γ 2), which may have implications in podocyte specific signaling and pathophysiology. This suggests that that new selective PPARy modulators can be potentially developed that will be able to distinguish between the two forms, $\gamma 1$ and $\gamma 2$, thus forming a basis of novel targeted therapeutic avenues.

Keywords: PPARy; podocyte; glomerular disease; proteinuria; alternative splicing

1. Introduction

Thiazolidinediones (TZDs) or peroxisome proliferator activated receptor γ (PPAR γ) agonists such as pioglitazone, directly protect podocytes from injury as demonstrated in podocyte culture in vitro studies [1–5], and reduce proteinuria and glomerular injury



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Copyright: © 2022 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). in various animal models of glomerular disease, as reported in preclinical in vivo studies [5–13]. Moreover, these beneficial effects of PPAR γ have been shown to be mediated by activation of podocyte PPAR γ as demonstrated by elegant studies using podocyte specific *Pparg* knock out (KO) mouse model [7,10]. This indicates a pivotal role for PPAR γ in maintaining glomerular function through preservation of podocytes [1,5–7,10]. Podocytes are highly differentiated epithelial cells in the kidneys, whose structural and functional integrity is critical for the maintenance of glomerular filtration barrier [14–16]. Accordingly, their dysfunction or loss is the initiating and progressing characteristic factor in a vast majority of renal diseases, leading to chronic kidney disease and kidney failure. Glomerular diseases characterized by high proteinuria manifest as nephrotic syndrome (NS) which is often associated with co-morbidities such as hypoalbuminemia, hypercholesterolemia, edema, and hyper-coagulopathy [17-23]. Recently, our group has further demonstrated that selective modulation of PPAR γ with a partial agonist, GQ-16, provides high efficacy in reducing proteinuria, as well as NS-associated comorbidities in an experimental model of nephrotic syndrome [24]. Thus, a multitude of evidence shows that targeting the podocyte PPAR γ pathway offers an attractive therapeutic strategy in NS. While PPAR γ has an established role in driving many adipogenic and lipid metabolizing genes mainly in adipose tissue, liver, and muscle, the understanding of molecular pathways regulated by PPAR γ in podocytes and glomeruli remains limited [5–7,9,10,25–29]. Furthermore, it is well-known that PPAR γ exists in two major isoforms, $\gamma 1$ and $\gamma 2$, which are a result of different promoter usage as well as alternative splicing (AS) [5,30,31]. As a result, $\gamma 2$ contains 30 additional amino acids at its N terminal, despite a longer mRNA sequence towards the 5'UTR of γ 1. While γ 1 is more widely expressed, γ 2 is mostly restricted to adipose tissue and liver. Few other forms have been discovered, including some with dominant negative function [32-37]. Moreover, cell-specific expression of PPAR γ variants has been demonstrated to play a differential role in downstream gene expression pattern in adipocytes (expressing the γ^2 variant) vs. the macrophages (expressing the γ^1 variant) [38]. However, the specific variant or isoform(s) expression of $PPAR\gamma$ and the role of its AS in podocytes and glomerular disease is unexplored. In this study, we sought out to answer this critical gap in the literature by hypothesizing that alternative splice variants of PPAR γ expressed in podocytes are distinct from adipocytes. To address this hypothesis, we performed a comprehensive analysis of different AS variants of PPARy in human podocytes and studied the expression of $\gamma 1$ vs. $\gamma 2$ variant in podocyte vs. adipose tissue in detail.

2. Materials and Methods

2.1. Podocyte Cell Culture and Treatments

Immortalized human podocytes were cultured in RMPI 1640 (Corning, Tewksbury, MA, USA) supplemented with 10% fetal bovine serum (FBS) (Corning, Tewksbury, MA, USA), 1% 100 X Penicillin Streptomycin L-Glutamine (PSG) (Corning, Tewksbury, MA, USA), and 1% Insulin-Transferrin-Selenium-Ethanolamine (ITS-X) (Gibco, Gaithersburg, MD, USA) [39]. Proliferating podocytes were cultured in a humidified atmosphere with 5% CO₂ at 33 °C and the media was changed twice per week and cells were passaged at ~70–80% confluence. Differentiation was induced by placing the cells at 37 °C under the same atmospheric conditions, for 14 days. Differentiated cells were treated with puromycin aminonucleoside (PAN) (Sigma-Aldrich, St. Louis, MO, USA) at 25 ug/mL or with pioglitazone (Alfa Aesar, Tewksbury, MA, USA) at 10 μ M. Control cells received treatment of vehicle dimethyl sulfoxide (DMSO) (Sigma-Aldrich, St. Louis, MO, USA). Cells were harvested after 24 h and total RNA isolated.

2.2. Kidney, Liver and Adipose Tissues

This study was approved by the Institution Animal Care and Use Committee at Nationwide Children's Hospital and the experiments were performed according to their guidelines. Male Wistar rats weighing ~150–200 g (Envigo, Indianapolis, IN, USA) were purchased, acclimated for 3 days, and housed under standard conditions. They were

sacrificed to collect kidneys, liver, and white adipose tissue (WAT) epididymal fat. Tissues were flash frozen in liquid nitrogen prior to RNA isolation. Glomeruli were isolated from harvested kidneys using the sequential sieving method as previously described [24] and total RNA was isolated. Commercially available human and rat kidneys total RNA was purchased from Zyagen (San Diego, CA, USA).

2.3. RNA Isolation

Rat epididymal fat and liver tissue samples were lysed in RNA extraction buffer (RLT) from the RNeasy kit (Qiagen, Germantown, MD, USA) with stainless-steel disruption beads for 4 min at 30.0 Hz using the Qiagen TissueLyser (Germantown, MD, USA). Total RNA was immediately isolated from the lysate using the RNeasy kit (Qiagen, Germantown, MD, USA), following the manufacturer's instructions. Total RNA was extracted from rat glomeruli tissue and human podocyte samples using the mirVanaTM Isolation Kit (Invitrogen, Carlsbad, CA, USA), according to the manufacturer's instructions. RNA yield and purity was checked prior to downstream applications by measuring the absorbance at 260, 280, and 230 nm with a nanodrop spectrophotometer (ThermoFisher, Waltham, MA, USA) and by calculating appropriate ratios (260/280, 260/230).

2.4. DNase Treatment, cDNA Synthesis, and PCR

Total RNA was DNase treated according to manufacturer's instructions (Zymo Research, Irvine, CA, USA). 500 ng-1 µg DNase-treated RNA was reverse transcribed using the iScript cDNA Synthesis Kit (Bio-Rad, Hercules, CA, USA), according to the manufacturer's instruction and the resulting cDNA was used for reverse transcription-polymerase chain reaction (RT-PCR) using HotStarTaq Plus Master Mix Kit (Qiagen, Germantown, MD, USA). The PCR conditions used were as follows: 95 °C for 5 min, 40 cycles [95 °C for 30 s, 50–60 °C (depending on the primer pair) for 30 s, 72 °C for 30 s], hold at 4 °C. Housekeeping gene *RPL6* was amplified only for 28 cycles to prevent over-saturation. The PCR-products were separated on a 1.5–2.5% agarose gel along with molecular weight ladder (ThermoFisher, Waltham, MA, USA), stained with 0.5 μ g/mL ethidium bromide (Research Products International, Mount Prospect, IL, USA), and images were captured using the Chemidoc (Bio-Rad, Hercules, CA, USA) equipment. Densitometry was performed using ImageJ Software (National Institutes of Health, Bethesda, MD, USA) and band density was calculated by subtracting background and normalizing to RPL6. Quantitative RT-PCR (qRT-PCR) was performed on samples using SYBR green (Bio-Rad, Hercules, CA, USA) on the Applied Biosystems 7500 Real-Time PCR System (Waltham, MA, USA). The PCR conditions were 95 °C for 10 min, 40 X (95 °C for 15 s, 50–60 °C for 1 min) followed by a melt curve to ensure specific products. Analysis was performed using the $\Delta\Delta$ Ct method [40] with normalization to RPL6.

2.5. Primer Design and Synthesis

Primers were custom designed to detect various PPAR γ variants for alternatively spliced mRNA products using the Reference Sequence Database (National Center for Biotechnology Information, Bethesda, MD, USA) and confirmed for the potential binding transcripts in the annotated database using the Primer Blast program (National Center for Biotechnology Information, Bethesda, MD, USA) (Table 1, Figure 1A). For variants with exon skipping, primers were designed to span the exon-exon junction to create a specific primer pair that would exclusively amplify the desired mRNA variant.



Figure 1. Alternatively Spliced Variants and Isoforms of PPAR γ in Podocytes. (A) The schematic depicts the exon and intron usage and the transcription and translation start and stop sites for various PPAR γ variants. The major isoforms of PPAR γ are isoforms $\gamma 1$ and $\gamma 2$, which encode for proteins containing 475 and 505 amino acids, respectively. The most common human annotated variants 1 (var1: NM_138712) and 2 (var2: NM_015869) encode isoforms y1 and y2 (depicted in the topmost PPAR γ illustration). These two variants are transcribed by different promoter usage by RNA Polymerase II (depicted in tall arrows) and alternative splicing (depicted by exon skipping). PPARy1 variant 1 includes exons A1 and A2, a start codon (ATG) on exon 1, and γ 2 variant 2 excludes exons A1 and A2, but includes exon B and a start codon in the same exon B. Thus, γ 1 protein product is 30 amino acids shorter than γ 2. The less known and understood variants are represented below, which include Variants 3, 4, 11, 9 and tORF4/Var 10. These are depicted with A1 and A2 usage in the schematic to emphasize those likely to be expressed in podocytes, although their adipocyte counterparts with exon B usage at the 5' UTR also exist. A1 and A2 exons are shown in light blue, exon B in dark blue, internal exons are in green boxes, introns are black lines, spliced out exons are marked with blue lines. Primer pairs used to amplify major variants and other variants are depicted in grey arrows and circles (A1, Ex1, a-j, Table 1). For the variants that skip an exon, the primer spans the exon-exon junctions and is shown as a split grey arrow underneath the introns (c, g, i). Protein isoforms are shown below their corresponding RNA in purple and amino acid (aa) length is listed. (B) Summary of the protein isoforms (γ) and RNA variants (var) with detailed exon/intron usage and annotations. γ^2 is found primarily in adipose tissue, while γ^1 is found in podocytes. Variants 1, 3, and 4 are variants of γ 1 that are likely to be found in podocytes, while variants 9, 10, and 11 are

additional less explored variants that would encode different isoforms. The variant 10 version of tORF4 is annotated that contains exon A1.2, but there is not an annotated version of tORf4 that would contain exon A1. (**C**) A representative gel showing the expression of several variants of PPAR γ in differentiated podocytes. PCR products were generated using specific primer pairs and run on an agarose gel and stained with ethidium bromide. All PPAR γ [primer pair Ex1 F, aR; 315 bp]; γ 2 [primer pair bF, aR; expected product 476 bp]; variants 1, 9, 11 (432 bp PCR product) and Var 4 (358 bp product) [doublet primer pair, A1F, aR]; variants 3, 5, and 10 (415 bp product); variant 7 (341 bp product); and variant 12 (503 bp product) [primer pair dF, aR]. Primers are listed in Table 1.

Name	Species	Forward	Reverse					
$PPAR\gamma$								
Total (All)	Homo Sapiens	ATGACCATGGTTGACACAG Ex1	GGAGTTGGAAGGCTCTTCAT a					
Doublet	Homo Sapiens	CGAGGACACCGGAGAGGG A1	GGAGTTGGAAGGCTCTTCAT a					
Var2 (γ2)	Homo Sapiens	TTTTAACGGATTGATCTTTTG b	GGAGTTGGAAGGCTCTTCAT a					
Var1 (γ1)	Homo Sapiens	GAAAGAAGCCAACACTAAACCAC c	GGAGTTGGAAGGCTCTTCAT a					
Var3 (γ1)	Homo Sapiens	TTCTGCTTAATTCCCTTTCC d	GGAGTTGGAAGGCTCTTCAT a					
Var4 (γ1)	Homo Sapiens	GCCGCCGTGGCCGCAGAAA e	GGAGTTGGAAGGCTCTTCAT a					
Var11 (γ3)	Homo Sapiens	CTTGCAGTGGGGATGTCTCA f	CAGCAAACCTGGGCGGTTGA g					
Var9 (γ5)	Homo Sapiens	AGAGCCTTCCAACTCCCTCA h	GTCATAGATAACGAATGGCAT i					
tORF4	Homo Sapiens	AGAGCCTTCCAACTCCCTCA h	AAACCCAAAACAACTTCCCG j					
Var1 (γ1) *	Rattus Norvegicus	GAAAGAAGCTGTGAACCACT Rat c	GGAGTTTGAAGGCTCTTCAT Rat a					
(γ2) *	Rattus Norvegicus	GCACTTCACAAGAAATTACC Rat b	GGAGTTTGAAGGCTCTTCAT Rat a					
Total (All) *	Rattus Norvegicus	ATTACCATGGTTGACACAG Rat Ex1	GGAGTTTGAAGGCTCTTCAT Rat a					
Podocyte Specific Genes								
NPHS1	Homo Sapiens	CGCAGGAGGAGGTGTCTTATTC	CGGGTTCCAGAGTGTCCAAG					
Nphs1	Rattus Norvegicus	TGCTCTTTGCAGTTGGTGGT	TCCTGATCCTGTCCTCCGAC					
SYNPO	Homo Sapiens	AGGTAGGCGTGGAGGAGG	GAGGTTCTGGTTGGGTTTGG					
Synpo	Rattus Norvegicus	CCGCTTGGGTCCCCTTC	TGAACTCGTTCACCCTCTGC					
WT1	Rattus Norvegicus/ Homo Sapiens	CCCTACAGCAGTGACAATTTATAC	TGCCCTTCTGTCCATTTC					
Adipose Tissue Specific Genes								
CD36	Homo Sapiens	GGACTGCAGTGTAGGACTTTCC	TTCCGGTCACAGCCCATTTT					
Cd36	Rattus Norvegicus	AAGTTATTGCGACATGATT	GATCCAAACACAGCATAGAT					
ADIPOQ	Homo Sapiens	ACTGCAGTCTGTGGTTCTGA	ACTCCGGTTTCACCGATGTC					
Adipoq	Rattus Norvegicus	TGTTCCTCTTAATCCTGCCCA	CCAACCTGCACAAGTTTCCTT					
ADIPSIN	Homo Sapiens	CGCCCCGTGGTCGGAT	GACAGCTGTAGCAGCAGGAG					
Adipsin	Rattus Norvegicus	AAGCTCTCCCACAATGCCTC	CATGGTACGTGCGCAGATTG					
AP2	Homo Sapiens	AAAGTCAAGAGCACCATAACCTT	TGACGCCTTTCATGACGCATTCC					
Ap2	Rattus Norvegicus	AAAGTGAAGAGCATCATAACCCT	TCACGCCTTTCATGACACATTCC					
Housekeeping Genes								
GAPDH	Homo Sapiens	AGACACCATGGGGAAGGTGA	GACGGTGCCATGGAATTTGC					
RPL6	Homo Sapiens	GCAACCCTGTCCTTGTCAGA	GCCAGCTGCTTCAGGAAAAC					
Rpl6	Rattus Norvegicus	AGGCATCAGGAGGGTGAGAT	TGTGAGGGTACATGCCGTTC					

Table 1. Primers Used in the Current Study.

* Rat nomenclature is annotated differently than human $\gamma 1$ and $\gamma 2$ variants. Primers _{a-j, Ex1, A1} are depicted in Figure 1A. Rat primers correspond to locations listed for human primers in Figure 1A.

2.6. PPAR-Promoter Responsive Element Prediction

Bulk RNASeq datasets from untreated control differentiated human immortalized podocytes (GEO GSE124622) [41] and untreated control differentiated human adipocytes (GEO GSE129153) [42] were analyzed. The average expression of control samples from the aforementioned GEO datasets were calculated and normalized expression greater than 2 were counted as detected. Genes were further cross referenced against the verified target and predicted PPAR Response Element (PPRE) genes that were downloaded from PPARgene database (http://www.ppargene.org/downloads.php (accessed on 31 July 2022)) [43] and using computational genomics approach [44]. Key podocyte and adipocyte specific genes measured in this study were also queried through the *PPAR* gene database (ppargene.org) to predict PPAR-responsive elements (PPRE) on their promoters Upon query submission, *p*-value and confidence level were generated if the gene was predicted as a PPAR target gene and contained putative PPREs in the 5 kb transcription start site flanking region. They were assigned into categories of confidence as: high (p > 0.8), medium $(0.8 \ge p > 0.6)$, and low $(0.6 \ge p > 0.45)$. A p value ≤ 0.45 was predicted as negative. Functional annotation of PPRE containing genes detectable exclusively in adipocytes vs. podocytes were performed using clusterProfiler (RRID:SCR_016884) [45] and plotted as many as 10 terms per cell type. The PPRE containing genes detectable in adipocyte vs. podocytes the overlapping genesets, and the associated functional terms were also compared.

2.7. Statistical Analysis

Statistical analysis was performed using GraphPad Prism Software version 8.2.0 for Windows (GraphPad Software, San Diego, CA, USA). Data are expressed as mean \pm standard error of mean and compared using unpaired Student's *t*-test. *p* value significance is depicted as * *p* < 0.05, ** *p* < 0.01, **** *p* < 0.0001.

3. Results

3.1. Podocyte Specific PPAR_γ Splice Variants

To identify the PPAR γ variants expressed in podocytes and to elucidate differences in podocytes vs. adipocytes, we examined the known variants of PPAR γ by reviewing the literature, scoping out the annotated versions (RefSeq database, NCBI), and by performing variant specific PCRs using custom-designed primers. A schematic of different variant forms (Figure 1A) and their exon/intron usage information and annotation (Figure 1B) is presented. Conducting PCR utilizing variant-specific primer pairs (Figure 1A and Table 1), we could identify total PPAR γ expression, specific variants corresponding to γ 1 (Var1 and 4) and other forms in differentiated podocytes, but not the $\gamma 2$ form (Var 2) (Figure 1C), which is highly expressed in differentiated adipocytes [46]. Variant 2 (encodes γ 2, a protein of 505 amino acids) includes exon B which contains a start codon. Variant 1 (encodes $\gamma 1$, a protein of 475 amino acids) utilizes exons A1 and A2 instead of B in the 5'UTR of the gene, resulting in the usage of a start codon ATG located on exon 1. Variants 3 and 4 both encode γ 1 as well, but they have differing 5'UTRs. Variant 3 contains an alternative exon A1.2, and variant 4 skips exon A2. Variant 11 has a similar 5'UTR to variant 1, but it skips exon 5, resulting in a frameshift and a premature stop codon in exon 6, encoding a potential product of 248 amino acids (γ 3). Variant 9 utilizes a start codon located in exon 3 and skips exon 4, encoding γ 5 (266 amino acids). Variant 10 or tORF4 uses a partially retained intron after exon 4 which contains a stop codon, making a product of 266 amino acids (γ 6). Variant 10 version of tORF4 is annotated and contains exon A1.2, but there is no annotated variant that uses exon A1. The possibility of expression of other variants was considered using primer pairs that would amplify specific products (Figure 1C). The doublet primer pair was able to discriminate between the cumulative expression of variants 1, 9 and 11 vs. variant 4. Cumulative expression of variants 3, 5, 7, 10 and 12 was ruled out using primer pair for Var3 (dF and aR), which would also amplify these other mentioned variants.

3.2. PPAR γ Splice Variants in Podocytes vs. Adipose Tissue PPAR γ

PPAR γ is a transcription factor and an established master regulator of fat cell physiology and differentiation in adipose tissue and liver [28,47]. While podocyte specific Pparg KO mice illustrated its role in podocytopathies [7,10], to understand the cell- and tissue-specific determinants of PPARy function in glomeruli and in podocytes, we performed specific PCRs to detect the expression of total (all) variants and exclusively of variant 1 (encoding γ 1) and variant 2 (encoding γ 2) in human podocytes and whole kidney and rat tissues of glomeruli, whole kidney, liver and WAT. In accordance with the hypothesis of cell specific action of PPAR γ , while expression of variant encoding the PPAR γ 2 isoform was observed in WAT, liver, and whole kidneys, $\gamma 2$ variant was undetected in human podocytes and in the rat glomeruli (Figure 2A). As shown in Figure 1A, γ 1 and γ 2 use different, promoter sites and undergo AS. Expression analysis of several uncommon variants described in Figure 1 demonstrated that they are present at low and varying levels in the whole human kidney (Figure 2B). Most of these (variants 4, 3, 11, 9, and tORF4) were found to be expressed at very low to undetectable levels in podocytes (Figure 2B). Moreover, analysis of the variants encoding γ 1 (Var 1 and 4) showed that their levels remained unchanged with PAN-induced injury of podocytes or treatment with PPARγ agonist, pioglitazone (Figure 2C).



Figure 2. Expression of PPARy Splice Variants in Podocytes vs. Adipose Tissue. (A) Total RNA was isolated from glomeruli, liver, and WAT samples harvested from rats, and from cultured differentiated human podocytes. Expression of the RNA variants encoding PPAR γ 1 and γ 2 (All variants), γ 1 only (Var1), and γ 2 only (Var 2) isoforms was measured by RT-PCR in isolated samples as well as commercially available human and rat whole kidney RNA. Rat total (all) γ [primer pair ex1 F, ex 2 R; 315 bp], rat y1 [primer pair exA2 F, ex2 R; 380 bp], rat y2 [primer pair exB F, ex2 R; 329 bp], rat Rpl6 [178 bp], human total (all) γ [primer pair Ex1 F, aR; 315 bp], human γ1 [primer pair cF, aR; 387 bp], human γ2 [primer pair bF, aR; 476 bp], Human RPL6 [370 bp]. (B) Several PPARγ mRNA AS variants were analyzed by RT-PCR in cultured differentiated human podocytes and commercially available human whole kidney RNA using the primer pairs drawn in Figure 1A and detailed in Supplementary Table S1. Var1 [primer pair cF, aR; 387 bp], Var3 [primer pair dF, aR; 415 bp], Var4 [primer pair eF, aR; 333 bp], Var11 [primer pair fF, gR; 242 bp], Var9 [primer pair hF, iR; 254 bp], tORF4 [primer pair hF, jR; 490 bp], Total [primer pair Ex1 F, aR; 315 bp], RPL6 [370 bp] (C) Expression of PPARγ mRNA variants 1 [primer pair cF, aR] and 4 [primer pair eF, aR], as well as total (All) PPARy [primer pair Ex1 F, aR] and GAPDH in puromycin aminonucleoside (PAN)-injured and pioglitazone-treated podocytes. Primers are listed in Table 1.

3.3. PPAR-Response Element (PPRE) Containing Genes in Podocyte vs. Adipocyte

PPAR γ binds to its promoter response elements (PPRE) along with other transcription factors such as retinoid X receptor (RXR) or to other DNA elements in association with factors such as nuclear factor kappa B (NF κ B), activator protein 1 (AP1), and in proximity to CCAAT enhancer-binding proteins (C/EBP α) [5,25,38]. We carried out promoter element prediction using a PPARgene database (ppargene.org (accessed on 31 July 2022)) [43] and cross-referenced it with bulk RNASeq GEO datasets obtained from control podocytes (GSE124622) [41] and adipocytes (GSE129153) [42]. We found that while 1669 PPREcontaining genes were detectable in both podocyte and adipocyte datasets, 453 were found to be unique to podocytes and 77 unique to adipocytes (Figure 3A(i), Supplementary Table S1). These were further classified into subsets based on the confidence level of PPRE prediction (see Methods; Figure 3A(ii)). We further analyzed a select set of PPRE containing genes known to play critical roles in podocyte pathophysiology, such as NPHS1 (nephrin), SYNPO (synpatopodin), and WT1 (wilms tumor 1) or those that have established roles in adipogenesis, such as CD36, ADIPOQ, ADIPSIN, and AP2 [29,47–52] (Table 2). All the adipogenic genes depicted high level of PPRE prediction, and NPHS1, SYNPO, and WT1 showed low to high levels of PPRE prediction. These genes which are known to play prominent roles in podocytes vs. adipocytes correlated with their expression in the GEO datasets in podocytes vs. adipocyte. In accordance, we also observed a marked disparity in the expression of these genes in podocytes vs. white adipose tissue (WAT) (Figure 3B). While the podocyte markers (NPHS1, SYNPO and WT1) are expressed at a higher level in podocytes vs. adipose tissue, expression of adipokines (ADIPOQ and ADIPSIN) and adipogenic genes (CD36 and AP2) was found to be high in WAT and absent or modestly expressed in podocytes (Figure 3B, Table 2). Furthermore, functional enrichment analysis of PPRE-containing genes exclusive to podocyte or adipocyte generated distinct maps of biological processes, cellular components, molecular functions and pathways (Figure 3C-F). While the adipocyte gene set was rich in genes involved in temperature homeostasis, lipid catabolic process, lipase activity and regulation of lipolysis, podocyte gene set was rich in genes involved in cell-cell adhesion and membrane transporter activity. Adipocyte PPREcontaining gene products were found located in lipid droplet and collagen-containing extracellular matrix, and podocyte PPRE-containing gene products were found in cell-cell junction and tight junction. Notably, functional enrichment analysis of all PPRE-containing genes from podocytes and adipocytes (including overlapping genes) generated mostly overlapping maps of biological processes, cellular components, molecular functions and pathways (Supplementary Figure S1A–D). We further cross-referenced the podocyte and adipocyte bulk RNASeq GEO datasets with PPRE genes from another independent source wherein PPREs were predicted in the conserved elements of within 5000 bps of transcription site of human genes [44]. We found that out of 1074 PPRE genes, 478 were detectable in both podocyte and adipocyte datasets, 139 were found to be unique to podocytes and 22 unique to adipocytes (Supplementary Figure S2A, Table S2). A total of 168 genes were common between the PPRE dataset and PPRE genes described by Lemay and Hwang [44] (Supplementary Figure S2B, Table S3). A majority of these overlapping genes were detected in both podocyte and adipocyte databases (Supplementary Figure S2C).

In summary, this data suggests the existence of a PPRE-containing gene expression, enrichment and pathway signature that is distinct in podocytes vs. adipocytes. This is likely driven by podocyte specific γ 1 form of PPAR γ vs. the adipocyte-specific γ 2 form.



Figure 3. PPAR_γ-Response Element (PPRE) Containing Genes in Podocytes vs. Adipocytes. (A) Normalized average log counts of untreated control human differentiated podocyte and adipocyte GEO datasets (GSE124622 and GSE129153, respectively) were cross-referenced against the verified target and predicted PPRE genes from the PPARgene database. (i) 1669 PPRE-containing genes were detectable in both podocyte and adipocyte datasets, 453 unique to podocytes and 77 unique to adipocytes. (ii) PPRE-containing genes classified by their confidence level. High-confidence (p > 0.8), median-confidence ($0.8 \ge p > 0.6$), low-confidence category ($0.6 \ge p > 0.45$). Genes with p value < 0.45 were predicted as negative. (B) Total RNA was isolated from cultured differentiated human podocytes and WAT harvested from rats. Expression of the podocyte marker genes (NPHS1 [Hu: 234 bp, Rat: 138 bp], SYNPO [Hu: 182 bp, Rat: 325 bp], and WT1 [Hu/Rat: 134 bp]) and adipose tissue genes (CD36 [Hu: 304 bp, Rat: 123 bp], ADIPOQ [Hu: 347 bp, Rat: 95 bp], ADIPSIN [Hu: 301 bp, Rat: 202 bp], and AP2 [Hu: 133 bp, Rat: 133 bp]), and housekeeping gene RPL6 [Hu: 370 bp, Rat: 178 bp] were analyzed. (i) Representative gels depicting the PCR products of key genes in podocytes and adipose tissue. (ii-iii) Quantitative mRNA fold-changes measured and graphically presented using RT-PCR and real time qRT-PCR quantification of key genes in podocytes and adipose tissue. Primers are listed in Table 1. (C-F) clusterProfiles generated functional enrichment of PPRE containing genes detectable exclusively in adipocytes (77) vs. podocytes (453) and plotted as 10 terms per cell type for (C) biological processes, BP, (D) cellular components, CC, (E) molecular functions, MF, and (F) kyoto encycopedia of genes and genomes, KEGG. The color of the dot indicates the intensity of adj p value (smaller adj p value is more red) and size of the dot indicates the proportion of genes from the term that are present in the cell-specific PPRE containing genes. * p < 0.05, ** p < 0.01, **** p < 0.0001.

Gene	Cells/Tissue Tested in the Current Study	<i>p</i> Value ^a	Confidence Level	PPREs Predicted ^b	Podocyte Count log ^c	Adipocyte Count log ^d (Normalized)
Podocyte Prominent Genes						
Nphs1	Podocyte/WAT	0.63696	Medium	4	2.27	-
Synpo	Podocyte/WAT	0.8715	High	5	7.89	9.03 (1.3)
Wt1	Podocyte/WAT	0.55556	Low	5	4.91	_
Adipocyte Prominent Genes						
Cd36	Podocyte/WAT	0.97886	High	3	3.77	846.26 (117.2)
Adipoq	Podocyte/WAT	0.97153	High	12	2.59	112.19 (15.5)
Adipsin	Podocyte/WAT	0.96216	High	6	4.22	991.32 (137.3)
Ap2/Fabp4	Podocyte/WAT	0.99997	High	10	_	1663.6 (230.5)
Housekeeping						
Rpl6	Podocyte/WAT	_	_	_	13.42	96.86 (13.42)

Table 2. Expression of Select PPRE Containing Genes.

^a Probability of PPAR target gene, higher value means a higher confidence. High-confidence (p > 0.8), medianconfidence ($0.8 \ge p > 0.6$), low-confidence category ($0.6 \ge p > 0.45$). Genes with p value ≤ 0.45 were predicted as negative. ^b Putative PPREs in the 5Kb transcription start site (TSS) flanking region. ^c Podocyte GEO Dataset # GSE124622. ^d Adipocyte GEO Dataset # GSE129153.

4. Discussion

PPAR γ agonism has a well-established beneficial role in the setting of diabetes, which has led to the generation of TZDs for the treatment of type II diabetes (Figure 4) [53,54]. Its beneficial role in kidney cells beyond its favorable systemic metabolic effects in diabetes originated from numerous preclinical studies and meta-analyses [55-59]. Earlier these effects of PPARy in influencing non-diabetic glomerular disease was thought to be through its anti-inflammatory actions on endothelial and myeloid cells [60-62]. However, recent discoveries and advances suggest the role of podocytes in mediating these effects. Seminal studies from our group and other research teams suggested that TZDs such as pioglitazone, directly protect podocytes from injury [1–5] and reduce proteinuria and glomerular injury in various animal models of glomerular disease [5–13] (Figure 4). Moreover, studies using podocyte specific *Pparg* KO mouse demonstrated that the beneficial effects of PPARγ are mediated by activation of podocyte PPAR γ , thus indicating a pivotal role for PPAR γ in maintaining glomerular function through preservation of podocytes [1,5–7,10]. However, the specific variant or isoform(s) expression of $PPAR\gamma$ and the role of its AS in podocytes and glomerular disease is unexplored. We hypothesized that AS variants of PPAR γ expressed in podocytes are distinct from adipocytes. We addressed this hypothesis by performing a comprehensive analysis of different AS variants of PPAR γ in human podocytes and by studying the expression of $\gamma 1$ vs. $\gamma 2$ variant in podocyte vs. adipose tissue in detail. Our findings suggest that the podocytes mainly express the y1 form, along with minimal expression of other AS forms. They do not express the γ^2 form, yet they are responsive to treatments by PPARy agonists, implicating that new selective modulators can be potentially developed that will be able to distinguish between the two forms, $\gamma 1$ and $\gamma 2$, thus forming a basis of novel targeted therapeutic avenues.



Figure 4. Illustration Describing the Roles of PPAR γ 1 and 2 Isoforms. Schematic of the suggested roles of PPARy 1 and 2 forms in regulating podocyte/glomerular disease specific effects (right panel) vs. metabolic/systemic and adipogenic effects (left panel). PPARy agonism plays a well-established role in the setting of treatment of Type II Diabetes, and treatment with its agonists, thiazolidinediones (TZDs), leads to increased ('a' pathway) insulin sensitivity and glycemic control [53,54]. However, these beneficial effects are also accompanied with adverse effects ('a' pathway), such as increased cholesterol, adipogenesis and decreased osteoblastogenesis [25]. TZDs or PPAR γ agonists have now been demonstrated to reduce podocyte injury ('b' pathway) by enhancing their viability and cytoskeletal integrity and decreasing apoptosis [1-5]. A few pathways involved in this process include crosstalk with the glucocorticoid receptor (GR), decreased caspase-3 activity and decreased TRPC6 expression. Gain of PPARy activity or its activation by TZDs ('c' pathway) has been shown to reduce proteinuria and glomerulosclerosis in various animal models of glomerular disease, such as 5/6 nephrectomy (Nx), minimal change disease (MCD), focal segmental glomerulosclerosis (FSGS) and glomerulonephritis (GN) [5–13,24]. On the other hand, loss of PPAR γ in podocyte specific *Pparg* KO mouse ('d' pathway) has demonstrated exacerbation of glomerular injury and proteinuria in animal disease models of FSGS and GN [1,5-7,10]. Analysis of the podocyte vs. adipose tissue specific expression of genes downstream of PPARy containing putative or identified PPREs (Figure 3) informs us that these downstream effects in podocytes/glomeruli are likely directed by the PPARy1 splice variant, distinct from the adipocyte-regulatory $\gamma 2$ variant.

Since their identification in 1990, PPAR γ has been recognized as a nuclear receptor superfamily member, a ligand-dependent transcription factor and a master regulator of adipogenesis and metabolism, which accounts for the insulin sensitizing effects of its agonists or anti-diabetic drugs such as TZDs [63]. While its role in regulating adipogenesis and lipid metabolism in adipose tissue, liver and skeletal muscle is well-characterized, the knowledge of PPAR γ -regulated signaling pathways in kidneys and in podocytes is scarce [5–10,25–29,64]. In our previous studies we speculated a cross-talk model between the PPAR γ and glucocorticoid receptor and further demonstrated that while pioglitazone could provide proteinuria reducing benefits, it could also enhance the ability of low-dose steroids to reduce proteinuria [6]. This suggested the potential clinical use of these FDA-

approved drugs in enabling the reduction of steroid dose, toxicity, and side effects to treat NS. We further translated these findings and demonstrated the ability of pioglitazone administration to improve clinical outcome in a child with steroid-refractory NS, which has now been extended in other NS patients as well [6,65]. Moreover, we have recently found that targeting PPAR γ with a selective modulator, GQ-16, offers an even better therapeutic strategy in NS, by reducing proteinuria and NS-associated side effects with a greater efficacy while providing reduced adipogenic effects than traditional TZDs [24]. However, to understand the proteinuria-reducing podocyte protective effects of PPAR γ activation, it is critical to build an understanding of podocyte specific transcriptional activity of PPAR γ as the lack of such knowledge is a barrier to modulating PPAR γ activity effectively. The existence of two isoforms $\gamma 1$ and $\gamma 2$ as a result of different promoter usage and AS further complicates the PPAR γ biology [5,30,31]. As a result, γ 2 contains 30 additional amino acids at its N terminal, despite a longer γ 1 mRNA sequence at its 5'UTR. We found that podocytes as well as glomeruli, express the more ubiquitous form of PPAR γ , the γ 1 form and not the γ 2 form, which is restricted to adipose tissue and liver. Interestingly a few other spliced forms of PPAR γ have been discovered in the past few years, which may confer varied functions [32–36]. Approximately 95% of mammalian genes undergo AS that can result in different protein products as well as microRNA sensitivity, mRNA stability, localization, and translation efficiency [66,67]. Moreover, AS is associated with many cellular processes and diseases such as differentiation, cancer, and immunity to name a few [66,68]. Moreover, expression of PPAR γ 1 and 2 mRNAs and isoforms in adipose tissue, liver, and skeletal muscle is known to be regulated in animal models of obesity, diabetes, and nutrition [69]. As the expression and role of the AS of PPAR γ in podocyte biology is unexplored, we performed a landscape analysis of other PPARy forms, which have been identified mostly in adipose tissues in recent years, including some with dominant negative function [32–37]. We could detect low expression of variants 4, 3, 11, and tORF4, and variant 9 was undetectable in the whole kidney. In podocytes, Var 9 was detectable at very low levels, and these additional variants were barely detectable consistently. We have recently also observed the presence of variant 11 form due to skipping of exon 5 (delta 5) in the WAT, which was particularly reduced with both pioglitazone and GQ-16 [24]. Ligand-mediated PPAR γ activation has been shown to induce skipping of exon 5 which by itself lacks ligand-dependent transactivation ability, and thus acts as a dominant negative form of PPAR γ [32]. In particular, recent evidence points towards distinct roles of y1 vs. y2 variant of PPARy governing specific and separate gene expression and metabolic functions at different stages in adipocytes [70], as well as between adipocytes and macrophages [38]. In the current study, analysis of the podocyte vs. adipocyte specific expression of genes downstream of PPARγ containing putative or identified PPREs informs us that these downstream effects in podocytes are likely directed by the PPAR γ 1 splice variant, distinct from the adipocyte-regulatory γ^2 variant (Figures 3 and 4). This is also evident from differential gene expression, enrichment and signaling pathways in the podocytes vs. adipocytes. Notably, we found podocyte specific PPRE-containing genes to be functionally enriched in cell-cell junction and cell-cell adhesion, which are known to play critical roles in the maintenance of podocyte structure and function and in its normal physiology [71]. This also suggests that activation of podocyte PPAR γ with agonists or selective modulators to drive these pathways is a viable therapeutic option to restore podocyte health during injury. At the same time, we acknowledge the limitations of in silico prediction of PPREs, species-specific differences that may exist in the location and functionality of these PPREs [72], and the involvement of other co-transcription factors that determine cell specificity of PPAR γ [38]. Overall, previous evidence indicates that the: (1) activation of PPAR γ results in reduction in podocyte injury and proteinuria [1,4–6,12,13], (2) podocyte specific PPAR γ KO results in resistance to protection of proteinuria by PPAR γ agonists [7,10], and (3) selective modulation of PPAR γ enhances proteinuria reducing efficacy with reduced adipogenic potential [24]. Taken together with the findings in the current study suggesting that the podocytes do not express the γ^2 form, yet they are

responsive to treatments by PPAR γ agonists, implicate that those new selective modulators can be potentially developed that will be able to distinguish between the two forms, γ 1 and γ 2, thus forming a basis of novel targeted therapeutic avenues.

PPAR γ also undergoes post-translational modifications and its differential phosphorylation at serine (Ser) 273 is an important determinant of its effect on adipogenesis and insulin sensitivity [47,73,74]. PPAR γ phosphorylation at Ser273 was demonstrated in obesity models and treatment with the PPAR γ agonist dephosphorylates this residue in the adipose tissue. GQ-16, a selective modulator of PPAR γ has been shown to dephosphorylate PPAR γ in an in vitro assay like traditional TZDs [73,74]. Since this site of phosphorylation on Ser273 is encoded by the last codon on exon 4, it remains conserved in major variants, 1 and 2, and several other variants in the current study, except for Var9, which was only detectable in podocytes at very low levels. It should be noted that the relative position is Ser243 in the variants including A1 and A2 exons instead of Ser273 in the B exon. Moreover, our recent findings suggested that relative phosphorylated PPAR γ levels (at Ser273) remained unaltered during glomerular injury and with treatments with pioglitazone and GQ-16 treatments in the glomeruli [24]. These data suggest that while PPAR γ -Ser273 is critical in determining the insulin sensitizing effects of PPAR γ in adipose tissue, it is unlikely to be a major mechanism of regulation in podocytes or glomeruli.

In the present study, we have observed a clear indication of the disparate roles of $\gamma 1$ vs. $\gamma 2$ in podocyte vs. adipocyte function and physiology and demonstrated the presence of other AS variants of PPAR γ mRNA in podocytes that would share the 5'UTR with variant 1 (encoding $\gamma 1$ isoform), although the roles of these other variants in podocytopathies and podocyte differentiation is still unclear. Their detection of differential expression is a challenge as they are observed to be present at too low levels in podocytes (unlike the adipocyte variants). It might be worthwhile to pursue studies in future wherein these variants are overexpressed using reporter plasmids followed by the analysis of their effects on podocyte physiology, actin cytoskeleton and function in the presence of injury causing agents, such as PAN. The demonstration of presence of these variants also warrants their detection in kidney biopsies from patients treated with PPAR γ agonists for diabetic nephropathy or type II diabetes.

In summary, we have shown that podocytes and glomeruli express the variants encoding the major form of PPAR γ 1 isoform and do not express the variants encoding the other major form γ 2. Moreover, expression of other AS forms of PPAR γ has been detected at very low levels in podocytes which may have implications in its pathophysiology. Future studies directed at developing new selective modulators that can distinguish between the two forms, γ 1 and γ 2, to enhance beneficial effects and reduce harmful effects of PPAR γ activation has the potential to provide novel targeted therapeutic avenues.

Supplementary Materials: The following supporting information can be downloaded at: https: //www.mdpi.com/article/10.3390/cells11213455/s1, Figure S1: clusterProfiles generated functional enrichment of all PPRE containing genes detectable in adipocytes and/or podocytes; Figure S2: PPAR γ -Response Element (PPRE) containing Genes in PPARgene dataset and Lemay and Hwang Manuscript detected in podocytes and/or adipocytes; Table S1: PPRE-driven genes from the PPARgene dataset, identified in podocyte and adipocyte datasets; Table S2: PPRE-driven genes from Lemay and Hwang manuscript, identified in podocyte and adipocyte datasets; Table S3: PPRE-driven genes in the PPARgene dataset and Lemay and Hwang manuscript

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