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Inhibition of Human *UGT2B7* Gene Expression in Transgenic Mice by the Constitutive

Androstane Receptor

MF Yueh, PL Mellon, and RH Tukey

Laboratory of Environmental Toxicology

Departments of Chemistry & Biochemistry, and Pharmacology -(MFY, RHT)

Department of Reproductive Medicine -(PLM)

University of California San Diego

9500 Gilman Drive MC 0722

La Jolla, CA 92093

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CAR and UGT2B7 Expression in Transgenic Mice

Address Correspondence to Robert H. Tukey

9500 Gilman Drive MC 0722

La Jolla, CA 92093

Phone-858-822-0286

Fax-858-822-0363

Email- rtukey@ucsd.edu

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Nonstandard Abbreviations:

UGT= UDP-glucuronosyltransferase

TCPOBOP= 1,4-b-s [2-(3, 5,-dichloropyridyloxy)]

CAR= Constitute androstane receptors

PXR= Pregnane X receptor

HNF= Hepatocyte nuclear factor

HDCA= Hydodeoxycholu acid

XenRs= Xenobiotic receptors

ABSTRACT

The x enobiotic r eceptors (XenRs), constitutive and r ostane r eceptor (CAR), and pr egnane X receptor (PXR) r equiate and all terit he metabolism of x enobiotic substrates. Among the 19 functional UDP- glucuronosyltransferases (UGTs) i n humans, UG T2B7 i s i nvolved in the metabolism of many structurally diverse xenobiotics and plays an important role in the clearance and detoxification of many therapeutic drugs. To examine if this gene is regulated by CAR and PXR in vivo, transgenic mice expressing the entire *UGT2B7* gene (*TgUGT2B7*) were created. Gene ex pression pr ofiles r evealed that *UGT2B7* is differentially ex pressed in liver, kidney, adipocytes, br ain, and es trogen-sensitive ti ssues, s uch a s ov ary and ut erus. Li ver UGT2B7 expression levels were decreased when TqUGT2B7 mice were treated with the CAR I igand 1, 4-b-s-[2-(3, 5,- dichloropyridyloxy)] (TCPOBOP), but not the P XR I igand pr egnenalone 16 αcarbonitrile. While TCPOBOP decreased the I evels of UG T2B7 mRNA in TgUGT2B7 mice it had no impact on $Tg(UGT2B7)Car^{-1}$ mice, adding support for a CAR- dependent mechanism contributing towards UGT2B7 gene suppression. Expression of promoter constructs in HepG2 cells showed the CAR dependent inhibition was linked to HNF4α-mediated transactivation of the *UGT2B7* promoter. The inhibitory effect of CAR on *UGT2B7* gene expression was validated in CHIP assays in which TCPOBOP treatment blocked HNF4α binding to the *UGT2B7* promoter. These r esults suggest that HNF 4α plays an important role in the constitutive expression of hepatic UG T2B7, and CAR a cts as a negati ve regulator by interfering with HNF 4α binding activity.

INTRODUCTION

Located in the cellular endople asmic reticulum, the fami by of UDP-glucuronosyltransferases (UGTs) plays a vital role in the metabolism and detoxification of numerous endogenous and exogenous compounds. There are 19 functional UGTs in humans, nine are encloded by the *UGT1* locus on chromosome 2, and the other *UGT2* genes on chromosome 4 (Mackenzie et al., 2005). The expression of these genes in human tissues is highly organized, with each tissue comprising its own complement of the UGTs (Gregory et al., 2004; Tukey and Strassburg, 2000). Among the human UGTs, UGT2B7 is expressed in many tissues and conveys broad substrates pecificity. Some estimates indicate that UGT2B7 is responsible for the metabolism of 35% of all clinical drugs (Williams et al., 2004). In addition, UGT2B7 participates in the metabolism of bile acids, fatty acids, and steroids (Ritter et al., 1992).

Since UG T2B7 p lays a k ey role in drug metabol ism and is abundant in human I iver (Izukawa et al., 2009) and intestine, efforts are underway to investigate the mechanisms leading to *UGT2B7* gene c ontrol. In human I iver, there e is large interindividual v ariability in the expression of UG T2B7 (Izukawa et al., 2009), part of which has been I inked to hepatocyte nuclear factor-1α (HNF1α) expression (Toide et al., 2002;Ormrod et al., 1999). In human Caco-2 cells, exposure to far nesoid X receptor (FXR) ligands, such as lithocholic acid, suppressed constitutive expression of UG T2B7 (Lu et al., 2005b). Retinoic acids, which are also metabolized by U GT2B7 (Samokyszyn et al., 2000) but play a keyrole in nuclear receptor function by activating the retinoid X receptor (RXR), have also been shown to suppress *UGT2B7* expression in Caco-2 cells (Lu et al., 2008). These results indicate that the family of xenobiotic nuclear receptors (XenRs), including FXR and possibly others that are expressed in liver and intestine such as the constitutive androstane receptor (CAR) and pregnane X receptor (PXR) may also be implicated in control of the *UGT2B7* gene.

The placement of human genes into mice that are expressed as transgenes serves as a powerful tool to ex amine the influence of hor mones, steroids and nuclear receptors towards influencing transcriptional control and function of the gene products. The generation of transgenic *UGT1* (*TgUGT1*) mice expressing the human *UGT1* locus has confirmed that the 9-*UGT1A* genes are expressed in a coordinated fashion (Chen et al., 2005) that resembles their expression pattern as mapped in human tissues (Strassburg et al., 1997a;Strassburg et al., 1997b;Tukey and Strassburg, 2000). The treatment of *TgUGT1* mice with ligands that activate the XenRs is a powerful tool to examine the role of these receptors in control and expression of the *UGT1A* genes, as the genes are regulated both through induction and tissue specificity (Chen et al., 2005;Ver reault et al., 2006;Yueh and Tukey, 2007;Senek eo-Effenberger et al., 2007). The functional role of the human *UGT1A1* gene in homeostatic control of serum bilirubin was recently demonstrated in humanized *UGT1* mice, which expresses the *UGT1A* genes in a complete *Ugt1*-null background (Fujiwara et al., 2010). We under took a similar approach to examine the regulation of the human *UGT2B7* gene.

The UGT2B7 gene spans 16 kbonchromosome 4 (Monaghan et al., 1994). We generated UGT2B7 transgenic mice (TgUGT2B7) with a bacterial a rtificial chromosome encoding the human UGT2B7 gene. Tissue-specific expression demonstrated by transcriptional levels revealed that the pattern of expression in TgUGT2B7 mice is comparable with what has been found for UGT2B7 expression in human tissues (Turgeon et al., 2001). Here we describe experiments which suggest functional inhibitory crosstalk between HNF 4α in liver of mice exposed to Tichonomy CPOBOP, confirming a right of the regulation of UGT2B7.

MATERIALS AND METHODS

Animals: The TgUGT2B7 mi ce were generated at the University of California San Diego Superfund Research Program Mouse Genetics Core Facility. A bacterial artificial chromosome (BAC) encoding the UGT2B7 gene (Genbank accession number RP13-644M16) was purified, microinjected into the pronucleus of CB6F 1 mouse eggs, and transplanted into the oviduct of pseudopregnant C57BL/6N m ice. For genotyping, DNA w as isolated from tail clippings, and a 418-bp DNA fragment in exon 1 or a 292-bp DNA fragment in Exon 6 were identified by PCR G ATTAAGAGATGGTCAGACC, Ex on 1 r (Exon 1 for ward: 5' everse: 5' CCACTTCTTCATGTCAAATATTTC; Exon 6 forward: AATTCAACATGATCAACCAGTG, Exon 6 reverse: GTCTCACCTATCAGGTTTTCC). Founders containing the UGT2B7 gene were bred with Car-null mice (Dr. M Negishi, NIEHS, Research Triangle Park, NC), and Tg(UGT2B7)Car^{+/-} mice bac kcrossed to prooduce $Tg(UGT2B7)Car^{-1}$ mice (genotyping for Car-null mice as described previously) (Ueda et al., 2002). All animals received food and w ater ad libitum and were housed in constant temperature rooms with a 12hr light/12 hr dark cycle. Mouse handling and experimental procedures were conducted in accordance with institutional guidelines.

UGT2B7 promoter activity: A 4-kb *UGT2B7* promoter element was cloned by PCR fr om the BAC DNA c ontaining the *UGT2B7* gene (Genbank ac cession number : PR13- 644M16) and subcloned into a pGL3 luciferase reporter plasmid. The primers for PCR-cloning of the *UGT2B7* promoter element were: -4kb (forward Kpnl: 5'ATTTGGTACCCAGTTCTCAGTA, reverse Bglll: 5' atttagatc ttcagtctgacac); - 2.8 k b (forward Kpn I: 5' atttggtac ctttgtgtgtcag,; r everse Bgl II: 5' aaaggaagatcttctatgggta); - 1.5kb (forward Kpn I: 5' taaaggtac caacagttcata, r everse Bgl II: 5' tgacagatcttgtttctgcag); - 0.4 k b (forward Kpn I: 5' attaggtac catgtttagtcatt, r everse Bgl II atttagatctggtgcaatgcaatg). Us ing the DNA fr agment s panning fr om –1.0 k b to the tr anslation start s ite (-1.0 k b for ward KpnI: 5' atttggtac ctaatgattaatgc, r everse XhoI: 5'

attactcgagacatcctggtgcaa), s ite-directed mutagenes is w as c arried out, al tering tw o bas es (underlined) on the HNF4 α core sequences (HNF4 α mutant, forward: 5' tatgtactttgcattataagggtt, reverse: 5' aacccttataatgcaaagtacata). For transient transfection experiments, HepG2 cells were seeded on 12- well pl ates 24 hr s befor e transfection. Cells were transfected with I uciferase plasmids allong with either pc DNA (Invitrogen, Carlsbad CA), HNF 4 α -pcDNA, or V P-CAR expression vectors (Xie et al., 2003) using lipofectamine 2000 (Invitrogen) based on the manufacturer's instructions. Cells were har vested with all ysis buffer (Promega, WI) 48 hours after the transfection, and the supernatant was collected by a brief centrifugation. The promoter activities were measured by the expression of firefly I uciferase and were normalized to the renilla luciferase levels using a dual luciferase reporter assay kit (Promega, WI).

Chromatin Immunoprecipitation (CHIP): CHIP anal ysis w as per formed us ing the mod ified protocol bas ed on the EZ -CHIP k it (Upstate Bi otechnology). HepG 2 c ells w ere tr ansfected either with an HNF4 α expression vector (HNF4 α -pcDNA) or an HNF4 α expression vector along with an activated CAR expression vector, V P-CAR (Xie et al., 2003). HepG 2 c ells w ere collected 48 hr s after the transfections and c ross-linked in DMEM (Invitrogen) containing 1% formaldehyde. The procedures for c ell I ysis and sonication to shear DNA were followed according to the manufacturer's protocol (EZ-CHIP kit, Upstate Biotechnology). One ml of cell extract in CHIP di lution buffer was pre-cleared by incubation with 60 μ I of Protein A Agarose/Salmon sperm DNA (Upstate Biotechnology) overnight at 4 °C. The cleared cellular extract was incubated with anti-HNF4 α antibody (Santa Crutz, CA) for 2 hr at 4 °C. Following precipitation with Protein A Agarose for 1h at 4 °C, the antibody-chromatin complex was then transferred to a spin column (Qiagen) for three 400 μ I washes with each of the following buffers: low-salt immune complex wash buffer, high-salt LiCI immune complex wash buffer (Okino et al., 2007), and Tris-

EDTA buffer. The protein-DNA complexes were eluted in 200 μ I elution buffer and DNA was then r everse c ross-linked and r eleased fr om the c omplex as i ndicated in the EZ -CHIP instructions. Following the DNA pur ification with spin columns, the pur ified DNA was further analyzed by real time PCR with a pair of primers (HNF4 α CH IP, for ward 5': gtgtgaacagttcatttaccttc; HNF4 α CHIP, reverse: 5' ctggtgcaatgcaatgctgt) for the amplification and quantification of the UGT2B7 promoter region containing the HNF4 α binding site.

Quantification of UGT2B7 gene transcripts by real time PCR: T otal RNA was i solated from tissues using Trizol (Invitrogen). One mi crogram of total RNA was used for the gener ation of cDNA with iScript cDNA synthesis kit (Bio-Rad, Hercules, CA). Following the cDNA synthesis, real time PCRs were conducted to determine Ct v alues using the MX 4000 Mul tiplex QPCR (Stratagene, La J olla CA). Br iefly, one mi crolitter of the cDNA templ ate from the RT -PCR reaction was used in a 20 µl of reaction mixture containing 10 µl of 2X MESA GREEN qPCR MasterMix (Eurogentec, San Di ego) and 0.4 µ M of a pair of primers for the detection of the mRNA of U GT2B7 or internal control gene cyclophilin (qPCR U GT2B7 for ward: 5' gacttttggttcgaaatatttgaca, qPCR UGT2B7 reverse: 5' gaggaaactgaaaattccagg; qPCR cyclophilin forward: 5' caga cgccactgtcgcttt, qP CR cyclophilin reverse: 5' tgtctttgg aactttgtctgcaa). The thermal profile is the following: 95°C for 10 min, 40 Cycles of 95°C for 40 sec, 58°C for 40 sec, and 72°C for 60 sec. After the amplification cycles were completed, the dissociation curve was generated at 95°C for 1 min followed by a 41 -dissociation cycle starting at 55°C and increasing by 1°C every 30 sec per cycle. Each sample was performed in triplicate and was quantified based on the formula Δ Ct = Ct (UGT2B7) — Ct (cyclophilin).

In vivo studies with TgUGT2B7 and Car-null mice: Age- matched groups of 8- 10 week old animals were used for all experiments. Wild type, Tg-UGT2B7, Car-- or Tg(UGT2B7)Car-- (n= 3

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or 4) mice were treated intraperitoneally every 24 hrs for 2 days with DMSO, PCN (10 mg/kg), dexamethasone (15 mg/kg), or TCPOBOP (4 mg/kg). All the chemicals were purchased from Sigma and di ssolved in 100 μ I DMSO for each injection. After 48 hr s, the I iver tissues, from each tr eatment gr oup, were pul verized in I iquid n itrogen and used for preparation of microsomes and total RNA. Microsomal fractions for UGT2B7 catalytic assay were prepared as described previously (Yueh et al., 2003).

Glucuronidation activity assay. UD P-glucuronyltransferase ac tivities were deter mined using HDCA and 4-hydroxyl estrone as substrates by TLC assay according to the method of Bans al and G essner with modification (Bansal and G essner, 1980). Br iefly, I iver tissues were homogenized in a five-fold volume of 1.15% ice-cold KC I and microsomal fractions were prepared in buffer (50 mM Tris-HCI (pH 7.6), and 10 mM MgCl₂) as described previously (Yueh et al., 2003). Each UGT assay was in a total volume of 100 μ I reaction mixture containing 50 mM Tris-HCI (pH 7.6), 10 mM MgCl₂, 100 μ M substrate, 500 μ M uridine 5-diphosphoglucuronic acid (UDPGA), 0.04 μ Ci of UDP[14 C]glucuronic acid, 8.5 mM s accharolactone and 75 μ g of microsomal protein. The reactions were performed at 37°C in a shaking water bath for 45 min. At the end of the reaction, 100 μ I of ethanol was added and the cell debris was pelleted by centrifugation. The supernatant was applied to TCL plates and chromatography performed in a mixture of (35:35:10:20 v/v) of n-butanol:acetone:acetic acid:water. The resulting glucuronides were visualized with a pho sphorimager (Molecular Dynamics Storm 820) and were removed and placed in scintillation fluid for quantification with a Liquid scintillation counter (Beckman Instruments, Palo Alto, CA).

Reagents: The BAC DNA containing UGT2B7 gene (PR13-644M16) was from Children's Hospital Oakland Research Institute (CHORI). 1,4-bis[2-(3,5-dichloropyridyloxy)]benzene

(TCPOBOP), pr egnenolone-16 α -carbonitrile (PCN), dex amethasone, a nd DMSO were fr om Sigma. Restriction enzymes and T4 DNA ligase for subcloning were from New England Biolabs. The Bradford assay for protein concentration analysis was from Bio-Rad. Taq polymerase, the dual-luciferase reporter assay system and reporter plasmids, pGL3-basic vector, pGL3 promoter vector and pRL- SV40 v ector were f rom Pr omega (Madison, WI). The expression v ector for HNF4 α (pc DNA-HNF4 α) was a k ind gift provided by Dr. Barbier at Lav al University Hospital Research Center, Q uebec, Canada. The construct for the expression vector VP- CAR was described previously (Xie et al., 2003). Thin-layer chromatography plates for the catalytic assay were from Whatman (Clifton, NJ).

RESULTS

Expression of UGT2B7 in transgenic mice

The organization of the *UGT2B7* ge ne in the BAC DNA, consisting of a 5' promoter region and 6 ex ons and introns, is shown in F ig 1. The BAC clone was purified and microinjected into fer tilized CB6F 1 mous e egg s, and *TgUGT2B7* transgenic mice were produced. The genotype analysis from tail DNA identified founders carrying sequences of exons 1 through 6. Three founders were used for breeding experiments to generate F1 progeny.

To determine if expression of the human gene in liver produced an intact mRNA, highly specific oligonucleotides were used to clone from reverse transcriptase product the ful I length UGT2B7 RNA i nto pc DNA followed by expression in COS-1 cells. Cell lysates prepared from UGT2B7 pc DNA transfected COS-1 cells displayed catalytic activity towards hyodeoxycholic acid (Fig 2A), a known substrate for UGT2B7. Enhanced levels of HDCA glucuronidation in liver microsomes from TgUGT2B7 mice when compared to wild-type mice where also observed (Figure 2B), confirming that expression of the UGT2B7 gene in transgenic mice produces a functional gene transcript.

Examination of the constitutive expression pattern of the *UGT2B7* gene was conducted by reverse transcription followed by PCR (RT-PCR) with UGT2B7 specific oligonucleotides to assess gene ex pression pr ofiles. The oligonucleotides used in the se experiments did not amplify gene transcripts from wild-type mouse liver RNA. Total RNA from different tissues was isolated from both male and female *TgUGT2B7* mice. The intense *UGT2B7* gene transcript was observed in liver and kidney tissues with liver being the most prominent (Fig 3). Lower levels of *UGT2B7* gene expression products were shown in large and small intestines, adipose tissue, brain, musicle, ov ary, and uter us. When we quantitated *UGT2B7* gene expression using real time-PCR procedures, the expression I evels matched the intensity of the banding patterns observed by RT-PCR. In experiments using human tissues, it has been demonstrated that the

UGT2B7 is expressed abundantly in various tissues including liver, kidney, small intestine, large intestine, mammary gland, and uterus (Ohno and Nakajin, 2009;Turgeon et al., 2001;Izukawa et al., 2009). O verall, the ti-ssue ex pression profile of the *UGT2B7* gene in transgenic mice corresponds well with that of humans indicating that the *TgUGT2B7* mice could be useful as an *in vivo* model to characterize *UGT2B7* gene expression.

Regulation of hepatic UGT2B7 expression by PXR or CAR ligands:

The effect of PX R and CAR activation of the *UGT2B7* gene in *TgUGT2B7* mice was evaluated following treatment with the P XR I igand PCN (10 mg/kg) or the CAR I igand TCPOBOP (4 mg/kg). Following a dministration by the input pour oute, Q-RT-PCR analysis to quantitate *UGT2B7* gene expression was conducted with RNA prepared from liver. PCN, a prototypical ligand of murine PXR, produced no effect on *UGT2B7* gene expression. However, treatment with TCPOBOP, a potent I igand of the mouse CAR, inhibited hepatic *UGT2B7* gene expression (Fig 4A).

In efforts to deter mine if CAR is tiled to riegulation of the *UGT2B7* gene, we crossed *TgUGT2B7* mice with *Car*^{-/-} mice to generate *Tg(UGT2B7)Car*^{-/-} mice. Wild- type, *TgUGT2B7*, or *Tg(UGT2B7)Car*^{-/-} mice were treated with either DMSO or TCPOBOP. RNA was prepared from liver t issues and the levels of *UGT2B7* gene ex pression quantitated by Q-RT-PCR. When compared with *TgUGT2B7* mice, the interruption of the *Car* gene in DMSO -treated *Tg(UGT2B7)Car*^{-/-} mice produced no change in *UGT2B7* gene expression (Fig 4B). TCPOBOP treatment to *TgUGT2B7* mice resulted in over an 80% reduction in gene expression. However, when *Tg(UGT2B7)Car*^{-/-} mice were treated with TCPOBOP, *UGT2B7* gene expression remained unchanged and were comparable to untreated mice. *Cyp2b10* gene expression, a well known TCPOBOP-inducible CAR target gene, were substantially increased by treatment of TCPOBOP in *TgUGT2B7* mice but not *Tg(UGT2B7)Car*^{-/-} mice (Fig 4C). Overall, these studies demonstrate that CAR functions as a negative regulator of the *UGT2B7* gene in liver.

Hepatocyte nuclear factor 4α (HNF4 α) is crucial for constitutive UGT2B7 expression in liver:

To study the mol ecular mechanisms that c ontrol constitutive expression of *UGT2B7* in liver, 4kb of the *UGT2B7* promoter was cloned from the BAC DNA and subsequently subcloned into a I uciferase reporter plasmid. HepG2 c ells were transfected with the *UGT2B7* promoter luciferase plasmids and high promoter activity was observed in the 400 bp proximal promoter region (-367/+12) adjacent to the transcription start site (Fig 5). Sequence analysis indicated there is one consensus DR1 core sequence (TGTACT X TGACTT) for HNF4α binding within this region. When HepG 2 cells were co-transfected with both a - 0.4 kb *UGT2B7* promoter-containing reporter plasmid (-0.4 kb/+ 0) and an HNF4α expression vector, the promoter activity was induced significantly, suggesting the presence of an HNF4α binding site in this region (Fig 5A). A two-base mutation in the DR1 core sequence blocked HNF4α-mediated transactivation, confirming the involvement of HNF4α in constitutive *UGT2B7* promoter activity (Fig 5B).

To explore the suppressive effect by CAR activation, HepG2 cells were transfected with an HNF4 α expression vector with or without cotransfection of a CAR expression vector (VPCAR). I nterestingly, transfection with HNF4 α alone increased promoter activity, and cotransfection of VP-CAR produced suppression of promoter activity (Fig 5C). Similar results were observed when HNF4 α transfected HepG2 cells were co-transfected with a CAR expression vector and treated with TCPOBOP for 48 hours, indicating that CAR might interact with HNF4 α and inhibit HNF4 α -directed transactivation. To gain further insight into the possible interaction of CAR and HNF4 α in regulating *UGT2B7* transcription, HepG2 cells were transfected with an HNF4 α expression vector with or without VP-CAR co-transfection followed by chromatic immunoprecipitation (CHIP) analysis. In CHIPs tudies using an HNF4 α anti-body, the precipitation of the DR1 element that contains the HNF4 α binding site (-181/+11), quantitated by real time PCR, was decreased in VP-CAR co-transfected HepG2 cells (Fig 6), indicating that the

inhibition of HNF4 α by CAR requires the inhibition in the binding of HNF4 α to the direct repeat 1

DISCUSSION

Recent studies have indicated that the UGT2B7 gene plays an important role in drug metabolism and s teroid homeostasis (Barbier et al., 2000; Coffman et al., 1998; Thibaudeau et al., 2006). The concern of species differences and lack of comprehensive knowledge regarding rodent UGT gene families prompted us to create a transgenic animal model containing a full length human UGT2B7 gene. The present study delineates the us e of this transgenic animal model to situdy their equilatory properties of the UGT2B7 gene. The expression pattern of UGT2B7 in various organs in TgUGT2B7 mice indicates that humor al and transcription factors meditating UGT2B7 gene ex pression r esemble thos e patter ns foun d i n human s. The observation that liver tissue had the highest expression levels of UGT2B7 suggested that liverspecific factors were required for physiological transcriptional responses. It has been shown that HNF4α pl ays an important role in regulating hepatic ex pression of phase II enzymes and transporters in mice (Lu et al., 2010). We provide evidence that HNF4α is the contributing factor responsible for constitutive expression of hepati c UG T2B7. HNF 4α regulates UGT2B7 gene expression by binding to a direct repeat motif of the AG GTCA's equence separated by one nucleotide (DR1) in the UGT2B7 5' fl anking pr omoter r egion. The H NF4α s pecificity and requirement for UGT2B7 gene ac tivation was further confirmed by mutation of the DR1 c ore sequence, which eliminated the b inding of HNF 4a to the promoter and abolished promoter activity. Si milar to our findings, mi ce I acking hepat ic HNF 4α had s ignificantly I ower gene expression of *Ugt2b1* when compared with wild type mice (Lu et al., 2010) indicating that both hepatic expressions of human *UGT2B7* and mouse *Ugt2b1* are controlled by HNF4α.

It is well documented that XenRs, PXR and CAR, act as xenobiotic sensors and mediate induction of numer ous x enobiotic metabolizing enz ymes. I nduction o f gl ucuronidation by xenobiotic receptors has been demonstrated using a number of clinical drugs and endogenous compounds. For example, CAR is a strong inducer of *UGT1A1* (Xie et al., 2003; Huang et al.,

2003), w hich proceeds through binding to a phenobar bital response element flanking the UGT1A1 gene pr omoter. We were surprised to observe that TCPOBOP treatment and activation of CAR in TqUGT2B7 mi celled to a reduction in UGT2B7 gene expression. The specificity of CAR-mediated regulation is supported by findings that PXR-specific ligands, such as PCN and dex amethasone, had no effect on the repression of UGT2B7 transcription in transgenic mice. Combined with evidence that overexpression of CAR produced a decrease in promoter activity of HNF 4α transactivation in HepG2 cells, UGT2B7 seems to be a c andidate gene for CAR- associated transcriptional inhibition. In addition, the role for HNF 4α in CARmediated inhibition of UGT2B7 expression was validated as CHIP as says revealed that CAR activation reduced HNF4α bound to the UGT2B7 chromatin. Activation of CAR inhibited HNF4α transactivation of UGT2B7 gene, which suggested that these two regulators are able to cross talk in the regulation of UGT2B7 expression. Finally, the us e of Car-null mice proved that the suppressive effect of T CPOBOP is linked to CAR, which acts as a transcriptional repressor in response to chemical activation by TCPOBOP and bl ocks HNF4α activation of UGT2B7 gene expression. By inhibiting HNF4α binding, CAR may prevent the changes in chromatin structure and consequent activation of UGT2B7 by HNF4α. In comparison with the antagonism between HNF4α and CAR for UGT2B7 gene regulation, a previous study showed that HNF4α inhibited PXR-mediated transactivation of CYP7A1 gene (Bhalla et al., 2004). The activated PXR did not affect the binding of HNF4α to CYP7A1. Instead, the association of HNF4α with cofactor PGC-1 (peroxisome proliferator activating receptor coactivator 1) bound to the promoter was inhibited. HNF4α-dependent transactivation of *UGT2B7* gene is mediated through the response element of the HNF 4α binding site in the promoter region, and a two-base change in the response element drastically reduces the ability of H NF4α to bi nd DNA. When acting as a po sitive regulator, CAR binds to the regulatory region of the target genes. Without a functional binding site in the UGT2B7 promoter region, CAR is able to interact with HNF 4α through a yet-to-be

identified mechanism that possibly involves contact with other associated transcription factors and cofactors which are specifically associated with the UGT2B7 promoter region. For example, CAR could be inhibitory by competing for binding to common coactivators for HNF4 α , such as PGC-1. Thus, the UGT2B7 specific regulation of HNF4 α and CAR may Targely depend on the promoter context.

UGT2B7 regulation at the transcription level is largely unstudied. Recently, using human Caco-2 c ells, UG T2B7 suppression by I ithocholic a cid w as I inked to n egative r egulation by farnesoid X receptor (FXR) (Lu et al., 2005a). Similarly, retinoids (i.e., all trans retinoic acid and 9-cis retinoic acid) were shown to inhibit UGT2B7 mRNA expression in this intestinal cell line. The fact that both lithocholic acid and retinoids are recognized as activators of CAR (Sakai et al., 2006; Chen et al., 2010) leads us to speculate that UG T2B7 down regulation in these human intestinal cells might be par tially claused by CAR activation. This downregulation of CARdependent UG T2B7 gene ex pression might have implications in metabolism of their apeutic agents des tined for glucuronidation by UG T2B7. Furthermore, CAR ac tivation may I ead to changes in the steady-state dynamics of steroids and bile acid homeostasis. A growing body of evidence shows the inhibitory effect of CAR on genes involved in hepatic glucose and lipid metabolism, bi le a cid b iosynthesis (Ueda et al ., 2002), such as phosphoenolpyruvate carboxykinase 1 (PEPCK1), gluc ose-6-phosphatase (G6P), and C YP7A1 activity (Miao et al., 2006). In combination, these studies implicate a diverse function of CAR as a negative regulator of genes as sociated with drug and xenobiotic, glucose, and I ipid metabolism. Compared with DMSO-treated Car- and w ild-type mi ce, T CPOBOP-treated Car- mice ex hibited hi gher UGT2B7 gene expression (~150%, Fig 7A); this finding is consistent with results from a number of other investigations (Bell and Michalopoulos, 2006; Tamasi et al., 2009) in which an induction of HNF4α by phenobarbital in the absence of CAR was observed. It is possible that TCPOBOP

is abl e to i influence HNF 4α activity in the absience of CAR, with induced levels of HNF 4α

contributing to greater *UGT2B7* gene expression.

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AUTHORSHIP CONTRIBUTIONS

Participated in research design: Yeuh, MF, Mellon, PL, and Tukey, RH

Conducted experiments: Yueh, MF

Contributed new reagents or analytical tools: Yueh, MF and Mellon PL

Performed data analysis: Yueh, MF

Wrote or contributed to the writing of the manuscript: Yeuh, MF and Tukey RH

REFERENCES

Bansal SK and Gessner T (1980) A Unified Method for the Assay of Uridine

Diphosphoglucuronyltransferase Activities Toward Various Aglycones Using Uridine

Diphospho[U-14C]Glucuronic Acid. *Anal Biochem* **109**:321-329.

Barbier O, Girard C, Breton R, Belanger A and Hum D W (2000) N-Glycosylation and Residue 96 Are Involved in the Functional Properties of UDP-Glucuronosyltransferase Enzymes.

Biochemistry 39:11540-11552.

Bell AW and Michalopoulos G K (2006) Phenobarbital Regulates Nuclear Expression of HNF-4alpha in Mouse and Rat Hepatocytes Independent of CAR and PXR. *Hepatology* **44**:186-194.

Bhalla S, Ozalp C, Fang S, Xiang L and Kemper J K (2004) Ligand-Activated Pregnane X Receptor Interferes With HNF-4 Signaling by Targeting a Common Coactivator PGC-1alpha. Functional Implications in Hepatic Cholesterol and Glucose Metabolism. *J Biol Chem* **279**:45139-45147.

Chen S, Beaton D, Nguyen N, Senekeo-Effenberger K, Brace-Sinnokrak E, Argikar U, Remmel R P, Trottier J, Barbier O, Ritter J K and Tukey R H (2005) Tissue-Specific, Inducible, and Hormonal Control of the Human UDP-Glucuronosyltransferase-1 (UGT1) Locus. *J Biol Chem* **280**:37547-37557.

Chen S, Wang K and Wan Y J (2010) Retinoids Activate RXR/CAR-Mediated Pathway and Induce CYP3A. *Biochem Pharmacol* **79**:270-276.

Coffman BL, King C D, Rios G R and Tephly T R (1998) The Glucuronidation of Opioids, Other Xenobiotics, and Androgens by Human UGT2B7Y(268) and UGT2B7H(268). *Drug Metab Dispos* **26**:73-77.

Fujiwara R, Nguyen N, Chen S and Tukey R H (2010) Developmental Hyperbilirubinemia and CNS Toxicity in Mice Humanized With the UDP Glucuronosyltransferase 1 (UGT1) Locus. *Proc Natl Acad Sci U S A* **107**:5024-5029.

Gregory PA, Lewinsky R H, Gardner-Stephen D A and Mackenzie P I (2004) Regulation of UDP Glucuronosyltransferases in the Gastrointestinal Tract. *Toxicol Appl Pharmacol* **199**:354-363.

Huang W, Zhang J, Chua S S, Qatanani M, Han Y, Granata R and Moore D D (2003) Induction of Bilirubin Clearance by the Constitutive Androstane Receptor (CAR). *Proc Natl Acad Sci U S A* **100**:4156-4161.

Izukawa T, Nakajima M, Fujiwara R, Yamanaka H, Fukami T, Takamiya M, Aoki Y, Ikushiro S, Sakaki T and Yokoi T (2009) Quantitative Analysis of UDP-Glucuronosyltransferase (UGT) 1A and UGT2B Expression Levels in Human Livers. *Drug Metab Dispos* **37**:1759-1768.

Lu H, Gonzalez F J and Klaassen C (2010) Alterations in Hepatic MRNA Expression of Phase II Enzymes and Xenobiotic Transporters After Targeted Disruption of Hepatocyte Nuclear Factor 4 Alpha. *Toxicol Sci* **118**:380-390.

Lu Y, Heydel J M, Li X, Bratton S, Lindblom T and Radominska-Pandya A (2005a) Lithocholic Acid Decreases Expression of UGT2B7 in Caco-2 Cells: A Potential Role for a Negative Farnesoid X Receptor Response Element. *Drug Metab Dispos* **33**:937-946.

Lu Y, Bratton S, Heydel J M and Radominska-Pandya A (2008) Effect of Retinoids on UDP-Glucuronosyltransferase 2B7 MRNA Expression in Caco-2 Cells. *Drug Metab Pharmacokinet* **23**:364-372.

Lu Y, Heydel J M, Li X, Bratton S, Lindblom T and Radominska-Pandya A (2005b) Lithocholic Acid Decreases Expression of UGT2B7 in Caco-2 Cells: a Potential Role for a Negative Farnesoid X Receptor Response Element. *Drug Metab Dispos* **33**:937-946.

Mackenzie PI, Walter B K, Burchell B, Guillemette C, Ikushiro S, Iyanagi T, Miners J O, Owens I S and Nebert D W (2005) Nomenclature Update for the Mammalian UDP Glycosyltransferase (UGT) Gene Superfamily. *Pharmacogenet Genomics* **15**:677-685.

Miao J, Fang S, Bae Y and Kemper J K (2006) Functional Inhibitory Cross-Talk Between Constitutive Androstane Receptor and Hepatic Nuclear Factor-4 in Hepatic Lipid/Glucose Metabolism Is Mediated by Competition for Binding to the DR1 Motif and to the Common Coactivators, GRIP-1 and PGC-1alpha. *J Biol Chem* **281**:14537-14546.

Monaghan G, Clarke D J, Povey S, See C G, Boxer M and Burchell B (1994) Isolation of a Human YAC Contig Encompassing a Cluster of *UGT2* Genes and Its Regional Localization to Chromosome 4q13. *Genomics* **23**:496-499.

Ohno S and Nakajin S (2009) Determination of MRNA Expression of Human UDP-Glucuronosyltransferases and Application for Localization in Various Human Tissues by Real-Time Reverse Transcriptase-Polymerase Chain Reaction. *Drug Metab Dispos* **37**:32-40.

Okino ST, Quattrochi L C, Pookot D, Iwahashi M and Dahiya R (2007) A Dioxin-Responsive Enhancer 3' of the Human CYP1A2 Gene. *Mol Pharmacol* **72**:1457-1465.

Ormrod D, Holm K, Goa K and Spencer C (1999) Epirubicin: a Review of Its Efficacy As Adjuvant Therapy and in the Treatment of Metastatic Disease in Breast Cancer. *Drugs Aging* **15**:389-416.

Ritter JK, Chen F, Sheen Y Y, Lubet R A and Owens I S (1992) Two Human Liver CDNAs Encode UDP-Glucuronosyltransferases With 2 Log Differences in Activity Toward Parallel Substrates Including Hyodeoxycholic Acid and Certain Estrogen Derivatives. *Biochemistry* **31**:3409-3414.

Sakai H, Iwata H, Kim E Y, Tsydenova O, Miyazaki N, Petrov E A, Batoev V B and Tanabe S (2006) Constitutive Androstane Receptor (CAR) As a Potential Sensing Biomarker of Persistent Organic Pollutants (POPs) in Aquatic Mammal: Molecular Characterization, Expression Level, and Ligand Profiling in Baikal Seal (Pusa Sibirica). *Toxicol Sci* **94**:57-70.

Samokyszyn VM, Gall W E, Zawada G, Freyaldenhoven M A, Chen G, Mackenzie P I, Tephly T R and Radominska-Pandya A (2000) 4-Hydroxyretinoic Acid, a Novel Substrate for Human Liver Microsomal UDP-Glucuronosyltransferase(s) and Recombinant UGT2B7. *J Biol Chem* **275**:6908-6914.

Senekeo-Effenberger K, Chen S, Brace-Sinnokrak E, Bonzo J A, Yueh M F, Argikar U, Kaeding J, Trottier J, Remmel R P, Ritter J K, Barbier O and Tukey R H (2007) Expression of the Human UGT1 Locus in Transgenic Mice by 4-Chloro-6-(2,3-Xylidino)-2-Pyrimidinylthioacetic Acid (WY-14643) and Implications on Drug Metabolism Through Peroxisome Proliferator-Activated Receptor {Alpha} Activation. *Drug Metab Dispos* **35**:419-427.

Strassburg CP, Manns MP and Tukey RH (1997a) Differential Down Regulation of the *UDP-Glucuronosyltransferase 1A* Locus Is an Early Event in Human Liver and Biliary Cancer. *Cancer Res* **57**:2979-2985.

Strassburg CP, Oldhafer K, Manns M P and Tukey R H (1997b) Differential Expression of the UGT1A Locus in Human Liver, Biliary, and Gastric Tissue: Identification of UGT1A7 and UGT1A10 Transcripts in Extrahepatic Tissue. *Mol Pharmacol* **52**:212-220.

Tamasi V, Juvan P, Beer M, Rozman D and Meyer U A (2009) Transcriptional Activation of PPARalpha by Phenobarbital in the Absence of CAR and PXR. *Mol Pharm* **6**:1573-1581.

Thibaudeau J, Lepine J, Tojcic J, Duguay Y, Pelletier G, Plante M, Brisson J, Tetu B, Jacob S, Perusse L, Belanger A and Guillemette C (2006) Characterization of Common UGT1A8, UGT1A9, and UGT2B7 Variants With Different Capacities to Inactivate Mutagenic 4-Hydroxylated Metabolites of Estradiol and Estrone. *Cancer Res* **66**:125-133.

Toide K, Takahashi Y, Yamazaki H, Terauchi Y, Fujii T, Parkinson A and Kamataki T (2002)
Hepatocyte Nuclear Factor-1alpha Is a Causal Factor Responsible for Interindividual
Differences in the Expression of UDP-Glucuronosyltransferase 2B7 MRNA in Human Livers.

Drug Metab Dispos 30:613-615.

Tukey RH and Strassburg C P (2000) Human UDP-Glucuronosyltransferases: Metabolism, Expression, and Disease. *Annu Rev Pharmacol Toxicol* **40**:581-616.

Turgeon D, Carrier J S, Levesque E, Hum D W and Belanger A (2001) Relative Enzymatic Activity, Protein Stability, and Tissue Distribution of Human Steroid-Metabolizing UGT2B Subfamily Members. *Endocrinology* **142**:778-787.

Ueda A, Hamadeh H K, Webb H K, Yamamoto Y, Sueyoshi T, Afshari C A, Lehmann J M and Negishi M (2002) Diverse Roles of the Nuclear Orphan Receptor CAR in Regulating Hepatic Genes in Response to Phenobarbital. *Mol Pharmacol* **61**:1-6.

Verreault M, Senekeo-Effenberger K, Trottier J, Bonzo J A, Belanger J, Kaeding J, Staels B, Caron P, Tukey R H and Barbier O (2006) The Liver X-Receptor Alpha Controls Hepatic Expression of the Human Bile Acid-Glucuronidating UGT1A3 Enzyme in Human Cells and Transgenic Mice. *Hepatology* **44**:368-378.

Williams JA, Hyland R, Jones B C, Smith D A, Hurst S, Goosen T C, Peterkin V, Koup J R and Ball S E (2004) Drug-Drug Interactions for UDP-Glucuronosyltransferase Substrates: A Pharmacokinetic Explanation for Typically Observed Low Exposure (AUCI/AUC) Ratios. *Drug Metab Dispos* **32**:1201-1208.

Xie W, Yeuh M F, Radominska-Pandya A, Saini S P S, Negishi Y, Bottroff B S, Cabrera G Y, Tukey R H and Evans R M (2003) Control of Steroid, Heme, and Carcinogen Metabolism by Nuclear Pregnane X Receptor and Constitutive Androstane Receptor. *Proc Natl Acad Sci USA* **100**:4150-4155.

Yueh MF, Huang Y H, Hiller A, Chen S J, Nguyen N and Tukey R H (2003) Involvement of the Xenobiotic Response Element (XRE) in Ah Receptor-Mediated Induction of Human UDP-Glucuronosyltransferase 1A1. *Journal of Biological Chemistry* **278**:15001-15006.

Yueh MF and Tukey R H (2007) Nrf2-Keap1 Signaling Pathway Regulates Human UGT1A1 Expression in Vitro and in Transgenic UGT1 Mice. *J Biol Chem* **282**:8749-8758.

FOOTNOTES

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LEGENDS FOR FIGURES

Fig 1. T he gene ar rangement of UGT2B7 in the BAC DNA: A 156 k b-bacterial ar tificial chromosome encoding the *UGT2B7* gene locus was used to generate the UGT2B7 transgenic

mice. The drawing is a representation of the *UGT2B7* gene locus, ranging from 48 kb to 64 kb,

in the BAC clone with 6 black boxes as 6 exons.

Fig 2. Deter mination of UG T ac tivity: (A) RNA fr om TqUGT2B7 mous e I iver w as i solated,

reverse transcribed to cDNA, subcloned into a pcDNA3 expression vector, and heterologously

expressed in COS-1 cells by transient transfections. Following preparation of cell lysates, UGT

activity was determined using HDCA as a substrate. UGT1A4 specific substrate amitriptyline

was as a negat ive control substrate. (B) UGT activity was determined in liver microsomes

prepared from TgUGT2B7 and wild type mice (n=4) using HDCA as substrate.

Fig 3. Tissue distribution of UGT2B7 transcript: Tissues from female and male TgUGT2B7 mice

were used to prepare total RNA. The UGT2B7 gene expression levels in various tissues were

examined by RT-PCR (female tissues) and real time-PCR using oligonucleotides specific for

UGT2B7 gene products and normalized to cyclophilin RNA. The specificity of the PCR product

was confirmed by direct sequence.

Fig 4. I nhibition of UG T2B7 expression by CAR I igand TCPOBOP and r eversion of UG T2B7

inhibition in Car null mice: (A) Age-matched TgUGT2B7 mice were treated with either DMSO,

CAR ligand TCPOBOP or PXR ligand PCN by i.p. injection for 48 hr s. The liver tissues were

used for preparation of total RNA. Following the reverse transcription for cDNA synthesis, real-

time PCR was conducted to determine Ct value with cyclophilin as an internal control gene. (B)

TgUGT2B7, Tg(UGT2B7)Car^{/-}, and wild type mice were treated with DMSO or TCPOBOP by i.p.

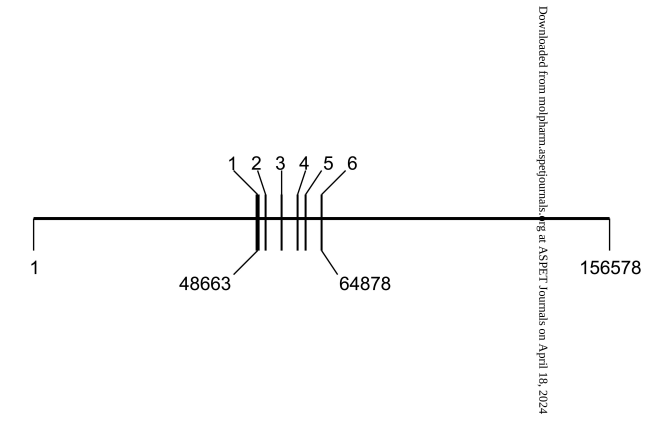
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injection for 48 hr s. RNA was isolated from the Liver tissues and the Levels of UG T2B7 mRNA were measured by real time PCR. (C) The Levels of mouse Cyp2b10 mRNA in Liver tissues of treated mice were examined by RT-PCR (Cyp2b10 forward: 5' aaagtcccgtggcaacttcc, Cyp2b10 reverse: 5' catcccaaagtctctcatgg).

Fig 5. T ransactivation of UG T2B7 promoter b v HN F4α and i nhibition of HNF 4α-mediated transactivation by CAR: A 4 kb of the UGT2B7 promoter was cloned, divided into 4 fragments, and subcloned into the luciferase reporter plasmids, pGL3 basic vector (BV) or promoter vector (PV). HepG 2 c ells w ere tr ansiently tr ansfected w ith UGT2B7 p romoter-containing r eporter plasmids, and I uciferase ac tivity w as deter mined in the c ytosolic f raction 48 hr s after transfections. (A) UGT2B7 promoter activities were compared bet ween cotransfection with a pcDNA plasmid or a HNF4α-containing expression vector and values were normalized to renilla luciferase activity by using a Luciferase dual a ssay kit (Promega) and were shown as fold induction. (B) Two bases were mutated, from AC to CA, in DR1-like core sequence within the UGT2B7 promoter region (UGT2B7 promoter - 1.0 k b/+0) by PCR-directed mutagenes is. The luciferase r eporter pl asmids c ontaining ei ther w ild t ype or mutated DR1 were tr ansiently transfected in to H epG2 ce lls. The promoter a ctivities w ere n ormalized t o r enilla lu ciferase activity and shown as firefly luciferase levels. (C) HepG2 cells were transfected with the reporter plasmid c ontaining the UGT2B7 promoter r egion (UGT2B7 pr omoter - 1.0 k b/+0) and cotransfected w ith HNF 4α, V P-CAR, or HNF4 α pl us VP- CAR. F orty eight hour s f ollowing transfection, fir efly lucifer ase activity was determined and values were normalized to renilla luciferase activity.

Fig 6. Chr omatic i mmunoprecipitation anal ysis of HN F4 α as sociated w ith the *UGT2B7* 5' flanking r egion. HepG 2 c ells w ere ei ther tr ansfected w ith a HNF 4 α expression v ector or cotransfected w ith an activated CAR expression v ector (VP-CAR). Transfected HepG 2 c ells were collected 48 hr after the transfections. Cells were fixed and sonicated for the preparation of sheared c hromatin, and i mmunoprecipitations w ere per formed us ing HNF 4 α anti body, or nonspecific IgG, as a negati ve c ontrol. Following i mmunoprecipitation, associated DNA w as amplified with a pair of primers targeting *UGT2B7* gene region -181 to +11, quantitated by real time PCR, and displayed by gel electorphorisis. Input and western blot of HNF4 α indicate equal amounts of lysates used prior to immunoprecipitation.

Fig 1



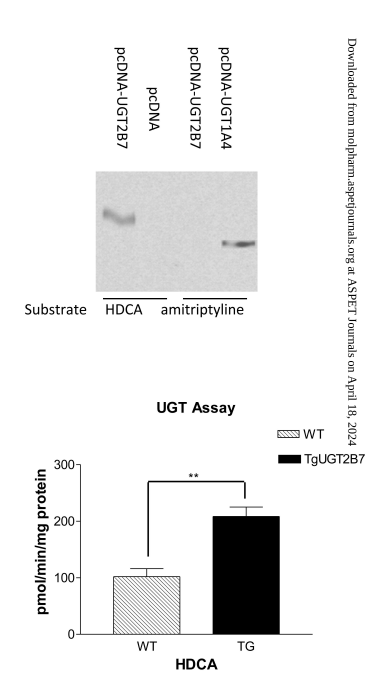


Fig 2B

Fig 2A

Fig 3

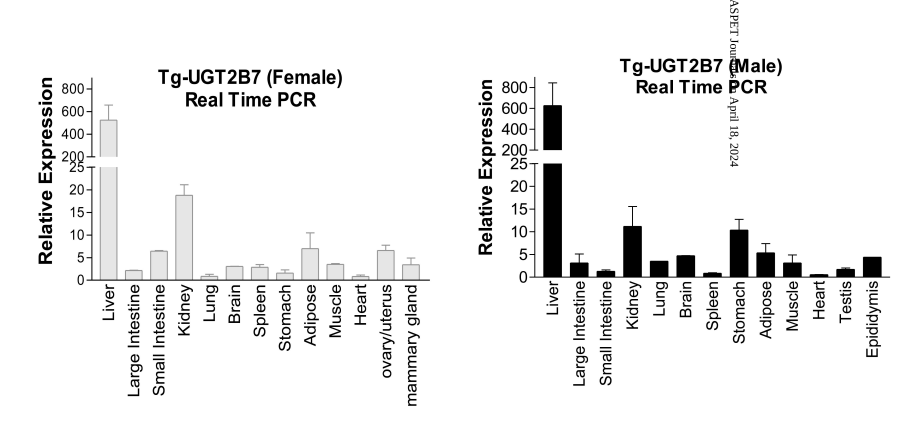
Tg-UGT2B7 (Female)

Tg-UGT2B7 (Female)

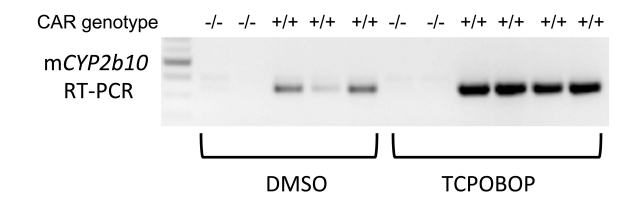
Tg-UGT2B7 (Female)

Tg-UGT2B7 (Female)

Real Time PCR



Downlook/Real Time PCR Fig 4A Fig 4B UGT2B7 mRNA/Real time PCR Relative Experssion 600-Relative Expression 500-400mRNA 300-200-0-Tg Tg/CAR^{-/-} Tg/CAR^{-/-} ASPET Journals on April 18, 2024 100-0-TCPOBOP PCN **DMSO** DMSO ТСРОВОР Fig 4C



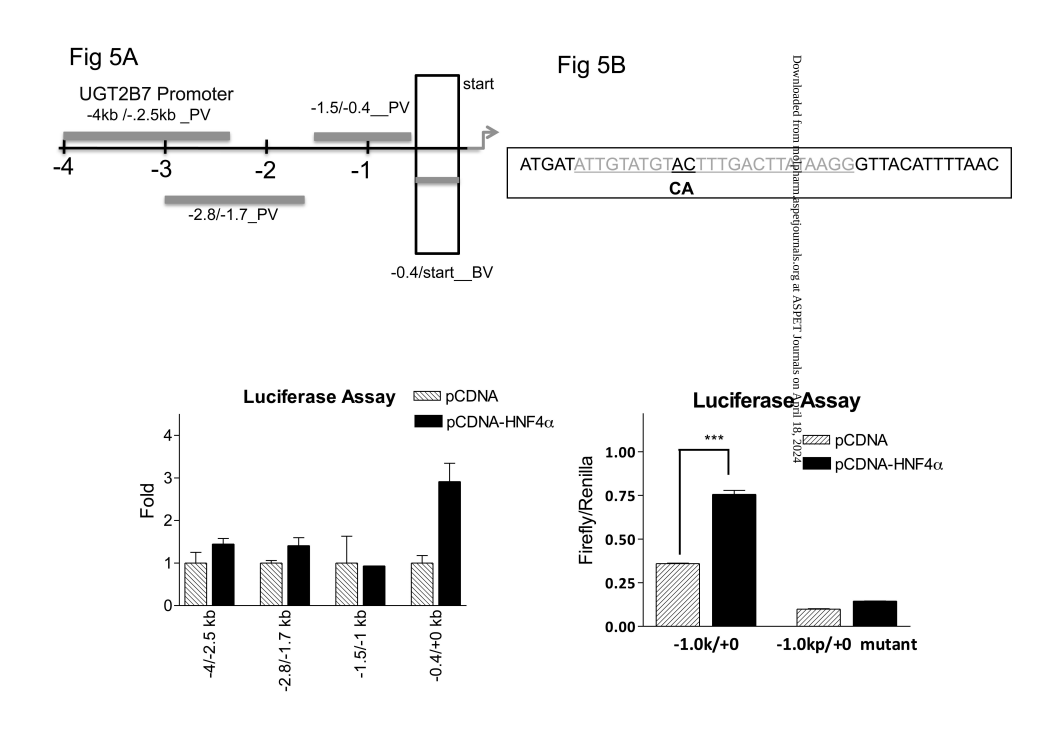


Fig 5C

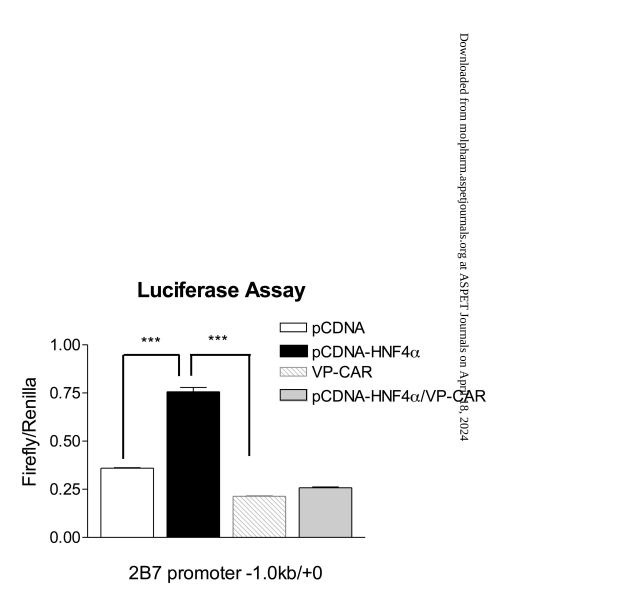


Fig 6

