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UGT Genomic Diversity: Beyond Gene Duplication

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1. Abstract

The human UDP-glucuronosyltransferase (UGT) superfamily comprises enzymes responsible for a major biotransformation phase II pathway: the glucuronidation process. The UGT enzymes are located in the endoplasmic reticulum of almost all tissues where they catalyze the inactivation of several endogenous and exogenous molecules including bilirubin, sex-steroids, numerous prescribed drugs and environmental toxins. This metabolic pathway is particularly variable. The influence of inheritable polymorphisms in human UGT-encoding genes has been extensively documented, and was shown to be responsible for a fraction of the observed phenotypic variability. Other key genomic processes are likely underlying this diversity; these may include copy-number variations (CNV), epigenetic factors and newly discovered splicing mechanisms. This review will discuss novel molecular aspects which may be determinant to UGT phenotypes.

Abbreviation: *UGT*, UDP-glucuronosyltransferase; *SNP*, Single nucleotide polymorphism; CNV; Copy-number variations, *AS*; Alternative splicing.

2. Introduction

UDP-glucuronosyltransferase (UGT) enzymes are key metabolic proteins that prevent accumulation of possibly toxic lipophilic compounds and initiate their elimination through more hydrophilic vehicules, such as biliary and renal systems (Dutton, 1980). This process is performed by the addition of a hydrophilic sugar moiety (glucuronide) from uridine diphosphate glucuronic acid (UDPGA) by UGTs (Dutton, 1980). The number of molecules handled by UGT enzymes is structurally divergent and almost infinite (Mackenzie, Bock et al., 2005; Mackenzie et al., 1997; Radominska-Pandya, Czernik, Little, Battaglia, & Mackenzie, 1999; Tukey & Strassburg, 2000b, 2001; Wells et al., 2004). This is related to the ability of these enzymes to catalyze the transfer of the glucuronic acid moiety at multiple functional sites, such as carbonyl, carboxyl, sulfuryl, hydroxyl, and amine groups (Mackenzie et al., 1997; Radominska-Pandya, Czernik, Little, Battaglia, & Mackenzie, 1999; Tukey & Strassburg, 2000b). The substrates for UGTs may be endogenous compounds, environmental products, or synthetic drugs. One may speculate that the vast substrate diversity of UGT enzymes would necessitate several enzyme-producing genes, and would be highly energy consuming for cells. However, higher eukaryotic organisms have developed several strategies allowing protein diversity without increasing the complexity of their genetic background. Based on current development in the field of glucuronidation and recent discoveries, we now better appreciate that the human *UGT* genes are certainly a remarkable example of this concept. Besides, this pathway is particularly variable and currently both genetic and environmental factors have been shown to determine individual glucuronidation phenotypes. In fact, the number of cases partially explained by genetic factors described

in the literature has increased exponentially and they were mainly focused on single-nucleotide polymorphism (SNP) (especially regulatory and missense mutations), insertions and deletions (indels) of nucleotides, for review see (Burchell & Coughtrie, 1997; Burchell et al., 2000; Guillemette, 2003; Strassburg, Kalthoff, & Ehmer, 2008), and more recently on copy number variation (CNV) (McCarroll et al., 2006). In this review, we will discuss novel molecular aspects which may be determinant to UGTs protein diversity and glucuronidation phenotypes.

3. The human UGT genes

The human UGT are a very good example of evolving protein diversity, mainly arising from either exon or gene duplication, but also by alternative promoter usage and splicing events (Mackenzie, Bock et al., 2005; Mackenzie et al., 1997; Owens, Basu, & Banerjee, 2005; Ritter et al., 1992). Indeed, the nature of the eukaryotic gene structure has evolved to facilitate protein multiplicity. This is possible by combining independent smaller parts or modules in varying contexts. Eukaryotic genes, in contrast to lower complexity organisms, are mostly characterized by small coding sequences (exons) intercepted with large non-coding genomic regions (introns) (Boeger et al., 2005). The junction of successive exons by the excision of intronic sequences in pre-mRNA molecules (mRNA splicing process) conducts to mature mRNAs, which may then be translated to polypeptidic products (Gilbert, 1978). Therefore, exon assembly, which determines protein domains, is the basis of the proteome diversity. Mechanisms inducing exon mixing, such as segmental duplication and recombination (exon shuffling) of genomic

regions, together with alternative pre-mRNA splicing events, are the major driving forces of such an evolving protein complexity (Moore & Proudfoot, 2009).

The UGT enzymes have been classified in two major families in human, namely *UGT1A* and *UGT2* (subdivided in *UGT2A* and *UGT2B*) (UDP Glucuronosyltransferase Homepage http://som.flinders.edu.au/FUSA/ClinPharm/UGT/index.html). The *UGT1A* family members are all encoded by a single gene locus on human chromosome 2q37.1, while several likely duplicated genes, located on 4q13.2, define UGT2B and UGT2A3 enzymes (Mackenzie, Bock et al., 2005). On the other hand, UGT2A1 and 2A2 proteins, similarly to *UGT1A*, arise from one locus on chromosome 4q13.3 (Mackenzie, Bock et al., 2005) (Court, Hazarika, Krishnaswamy, Finel, & Williams, 2008).

3.1 The human UGT1A family members

The human *UGT1A* gene well exemplifies the usage of alternative promoter (consequently the first exon) to increase protein diversity. The complexity of the *UGT1A* locus has been firstly highlighted by the discovery of bilirubin transferase cDNAs and phenol transferase cDNAs having unique 5' sequences but identical 3' ends (Gong et al., 2001; Owens, Basu, & Banerjee, 2005; Ritter et al., 1992). Further experiments clearly indicated that these diverging cDNAs are encoded by one gene locus, and lead to transferases having various, sometimes overlapping and divergent substrate specificities. In fact, the human *UGT1A* locus is defined by 13 first exons, which are alternatively spliced to four (more recently five) common exons (**Figure 1**). From the 13 possible mRNA isoforms, nine conduct to functionally active polypeptides (UGT1A1, 1A3, 1A4, 1A5, 1A6, 1A7, 1A8, 1A9 and 1A10) and 4 pseudogenes (UGT1A2p, UGT1A11p and

UGT1A12p, UGT1A13p) (Gong et al., 2001; Ritter et al., 1992). The first exons most probably encode for the substrate-binding domain (Meech & Mackenzie, 1997a). Based upon their function and sequence similarity, UGT1A members are divided in two groups. the bilirubin-like-associated enzymes, which are defined by UGT1A1, UGT1A3, UGT1A4 and UGT1A5, and the phenol-like-associated group, encompassing UGT1A6, UGT1A7, UGT1A8, UGT1A9, and UGT1A10 (Owens, Basu, & Banerjee, 2005; Ritter et al., 1992). This family of enzymes has the ability to conjugate several xenobiotics. such as drugs and carcinogens (for a more detailed review see (Court, 2005; Dutton, 1980; Guillemette, 2003; Mackenzie, Bock et al., 2005; Mackenzie et al., 1997; Owens, Basu, & Banerjee, 2005; Ritter, 2000; Tukey & Strassburg, 2000a)), but are also involved in metabolism of endogenous compound, namely bilirubin (Ritter et al., 1992; Ritter, Crawford, & Owens, 1991), bile acids (Gall et al., 1999b; Ritter, Chen, Sheen, Lubet, & Owens, 1992; Trottier et al., 2006), serotonin (Court, 2005), androgenic and estrogenic steroid hormones (Belanger et al., 1998; Belanger, Pelletier, Labrie, Barbier, & Chouinard, 2003; Guillemette, Belanger, & Lepine, 2004; Lepine et al., 2004; Levesque et al., 1997; Starlard-Davenport, Lyn-Cook, & Radominska-Pandya, 2008). Nucleic acid sequence similarity clearly argues for ancestral exon duplication among UGT1A members. In addition, an inter-species sequence comparison of UGT1A first exons demonstrated that UGT1A1 and UGT1A6 are ancestrally common exons, and that other isoforms arose more recently in the evolution (Owens, Basu, & Banerjee, 2005).

A number of genetic polymorphisms, especially non-synonymous mutations, have been reported in UGT1A first exons (for a comprehensive list of UGT polymorphisms: http://www.ugtalleles.ulaval.ca). Considering the role of exon 1-derived UGT1A peptides

in substrate binding and the continuously changing environment, these coding polymorphisms indicate that UGT1A is an evolving locus armed to cope for new encountered exogenous components. In fact, some mutations are exclusively or mostly observed in particular human populations, supporting an environmental pressure in sequence selection. This is well exemplified by UGT1A1 mutations, UGT1A1*6, UGT1A1*7, and UGT1A1*27 solely detected in Asians (Aono et al., 1995; Aono et al., 1993). Moreover, examples of SNP association with glucuronidation rates and substrate selectivity support this evolutionary mechanism. A comparative study between UGT1 loci from diverse vertebrates indicates that exon birth-and-death is likely involved in UGT evolution, as demonstrated by pseudogenes (Riedy et al., 2000; Turgeon, Carrier, Chouinard, & Belanger, 2003). The shared and also more conserved region of the protein (for UGT1A1 this corresponds to amino acids 289 to 533) includes the transmembrane domain, the binding domain for the co-substrate UDPGA (Meech & Mackenzie, 1997a). This conserved proteins domains are essential to preserve the fundamental glucuronic acid transferase function (Ouzzine, Barre, Netter, Magdalou, & Fournel-Gigleux, 2003; Patana, Kurkela, Goldman, & Finel, 2007; Radominska-Pandya, Czernik, Little, Battaglia, & Mackenzie, 1999).

The expression of each UGT1A mRNA isoform is determined by their own promoter region (Gong et al., 2001; Ritter et al., 1992). The use of multiple promoters by UGT1A gene provides a genomic framework enabling a cell to respond to diverse physiological and pathophysiological stimuli. Consequently, UGT1A enzymes also diverge in their expression profile in tissues and between individuals (Burchell & Coughtrie, 1997;

Guillemette, 2003; King, Rios, Green, & Tephly, 2000; Mackenzie, Gregory et al., 2005; Strassburg, Kalthoff, & Ehmer, 2008; Strassburg, Nguyen, Manns, & Tukey, 1998; Tukey & Strassburg, 2000b). For instance, while UGT1As are mostly produced in gastrointestinal tissues (GI) (eg. liver, stomach, small intestine, and colon), some are specifically encountered in extrahepatic tissues (UGT1A7, UGT1A8 and UGT1A10). As a major contributor in liver's detoxification process, UGT1A1 is widely expressed throughout human body, whereas UGT1A9 is largely expressed in liver and kidney (Ritter, 2007; Strassburg, Manns, & Tukey, 1998; Strassburg, Nguyen, Manns, & Tukey, 1998; Tukey & Strassburg, 2000a). UGT1A6, UGT1A3 and UGT1A4 are also produced in liver, and along with UGT1A10, in GI tract. Some UGTs are also present in brain, ovary, uterus, prostate and breast (Belanger, Pelletier, Labrie, Barbier, & Chouinard, 2003; Guillemette, Belanger, & Lepine, 2004; King, Rios, Green, & Tephly, 2000; Starlard-Davenport, Lyn-Cook, & Radominska-Pandya, 2008; Strassburg, Kalthoff, & Ehmer, 2008). Thus, it is clear that the alternative usage of duplicated first exons is the basis of UGT1A function and expression diversities. Besides, all this tissue-specific expression is well orchestrated by the cooperative interaction between transcription factors and the transcription initiation complex (for review see (Mackenzie et al., 2003; Owens, Basu, & Banerjee, 2005; Strassburg, Kalthoff, & Ehmer, 2008; Tukey & Strassburg, 2001) (Gardner-Stephen & Mackenzie, 2008) Studies concerning UGT1A promoter-interacting factors have revealed the binding of some important activators, such as HNF-1, cdx2, Nrf2, PXR, XRE (AhR) and CAR (Aueviriyavit, Furihata, Morimoto, Kobayashi, & Chiba, 2007; Bonzo, Belanger, & Tukey, 2007; Buckley & Klaassen, 2009; Gregory, Lewinsky, Gardner-Stephen, & Mackenzie, 2004; Senekeo-Effenberger et al., 2007; Sugatani, Sueyoshi, Negishi, & Miwa, 2005; Verreault et al., 2006; Yuan, Li, & Yang, 2007). Transcriptional regulation of the *UGT1A* gene by transcription factors is more specifically addressed in some following chapters. Variability in these regulatory mechanisms are also accounting for interindividual differences in the glucuronidation pathway (Aueviriyavit, Furihata, Morimoto, Kobayashi, & Chiba, 2007; Mackenzie, Gregory et al., 2005; Ramirez et al., 2008; Toide et al., 2002). In addition, the recent discovery of a new alternatively used exon (referred as exon 5b) in the common region of the locus (exon 2-5) sheds light on innovative mechanisms likely implicated in the control UGT1A-mediated glucuronidation (see next section) (Girard et al., 2007; Levesque, Girard, Journault, Lepine, & Guillemette, 2007).

3.2 The human UGT2 family members

In contrast to the *UGT1A* locus, the human *UGT2* family members are encoded for by unique and multiples genes on 4q13 (Mackenzie, Bock et al., 2005). As illustrated in **Figure 2**, *UGT2B* and *UGT2A* genes likely originated from segmental chromosome duplication. Indeed, the phylogenetic relationship well demonstrates the prominent similarity between these genes. *UGT2A3* and all *UGT2B* genes are composed of at least six coding exons (Riedy et al., 2000; Turgeon et al., 2000) (Court, Hazarika, Krishnaswamy, Finel, & Williams, 2008). UGT2A1 and UGT2A2 mRNAs are transcribed by the same locus, by the use of alternative first exons, as demonstrated for the *UGT1A* locus (Jedlitschky, Cassidy, Sales, Pratt, & Burchell, 1999; Mackenzie, Bock et al., 2005). The intergenic distances vary greatly and the gene transcription may occur on either strand. As pictured in **Figure 2**, and similar to the *UGT1A* family, the first exon

encodes for the substrate-binding domain, and the other coding exons are responsible for the UDPGA-binding and transmembrane domains. One of the most studied UGT2B enzyme is UGT2B7. Of great importance in the liver, it is abundantly expressed in GI tract, but also in breast, uterus, kidney, brain, and other tissues (Turgeon, Carrier, Levesque, Hum, & Belanger, 2001). UGT2B7 has the ability to glucuronidate several endogenous compounds, such as steroid hormones (Gall et al., 1999a), retinoids (Czernik, Little, Barone, Raufman, & Radominska-Pandya, 2000), fatty acids (Jude, Little, Bull, Podgorski, & Radominska-Pandya, 2001), but also exogenous molecules such as morphine (Coffman, Rios, King, & Tephly, 1997), AZT (Barbier et al., 2000), EFV (Mutlib et al., 1999), NSAIDS (C. Jin, J. O. Miners, K. J. Lillywhite, & P. I. Mackenzie, 1993; C. J. Jin, J. O. Miners, K. J. Lillywhite, & P. I. Mackenzie, 1993), and others (Court, 2005). Although the other UGT2B family members have been investigated to a smaller extent, they have important role, especially with endogenous compounds. For example, UGT2B4 is involved in the metabolism of catechol estrogens and bile acids (Fournel-Gigleux, Jackson, Wooster, & Burchell, 1989; Levesque, Beaulieu, Hum, & Belanger, 1999; Pillot et al., 1993; Ritter, Chen, Sheen, Lubet, & Owens, 1992). UGT2B15, UGT2B17 and UGT2B28 arise as the major UGT2B sex-steroid metabolizing transferases (Beaulieu, Levesque, Hum, & Belanger, 1996; Levesque et al., 1997; Levesque et al., 2001). In contrast to the UGT1A first exons, it appears that fewer mutations affecting the coding regions occurred in these genes, which indicates a more conservative structure for *UGT2B* genes [48-50]. Polymorphic deletions were shown to occur for UGT2B genes namely UGT2B17 and UGT2B28 (Jakobsson et al., 2006; McCarroll et al., 2006; Wilson et al., 2004). In fact, copy-number variations of these two genes have been described with divergent frequencies among ethnic populations and surely contribute as a source of phenotypic differences (Spielman et al., 2007). These will be further discussed (see section 7).

4. Novel UGT-associated mRNA isoforms

With the recent genome-wide analyses, the alternative splicing becomes an important mechanism explaining the transcriptome (which is defined as all "transcripts" in a cell type at a given time) diversity, with the vast majority of human genes having alternatively spliced mRNA isoforms (Johnson et al., 2003; Kampa et al., 2004; Maniatis & Tasic, 2002; Stamm, 2008; E. T. Wang et al., 2008). There are several ways to generate alternative mRNA isoforms from one gene locus, namely the alternative promoter usage (observed in *UGT1A* gene), alternative exon usage, intron retention and the alternative 3'UTR (**Figure 2**) (Kornblihtt, 2005; Moore & Proudfoot, 2009). Beside to dramatically increases the transcriptome diversity, alternative splicing introduces novel mRNA molecules, which may lead to divergent polypeptides, in term of biological function and/or expression profile, but also having intrinsic regulatory role. Recent evidence support that this is the case for human *UGT* genes (Girard et al., 2007; Levesque, Girard, Journault, Lepine, & Guillemette, 2007).

The human genome project has provided to the scientific community priceless data about genomic organization and sequences, but has also contributed to extend knowledge on gene products, especially RNA species. In fact, sequencing of large cDNA libraries (expressed-sequence tag, EST), the use of Tag-based sequencing methods such as SAGE (Anisimov, 2008)and CAGE (de Hoon & Hayashizaki, 2008), the development of

specialized expression arrays (exon and tiling arrays) (Samanta, Tongprasit, & Stolc, 2007) (Lin et al., 2009), and the use of "next-generation" sequencing system to perform RNA-seq (Z. Wang, Gerstein, & Snyder, 2009), are all important tools to better understand the transcriptome.

As abovementioned, the UGT1A locus is a perfect example of "one locus given several divergent mRNAs". The recent discovery of a new alternative exon in the common region of the *UGT1A* locus further supports such an increased mRNA diversity (Girard et al., 2007; Levesque, Girard, Journault, Lepine, & Guillemette, 2007). Considering the high occurrence of gene-associated mRNA diversity, it is likely to investigate whether other *UGT* loci have additional mRNA species. In the following, we listed evidence for additional UGT mRNA isoforms by analyzing expressed-sequence tags (EST) gene clusters. We also mentioned some experimental supports for such mRNA species.

4.1 UGT-derived EST clusters

UniGene (http://www.ncbi.nlm.nih.gov/unigene) is an NCBI database attempting to clusterize transcribed sequences according to their similarity, thus providing a set of transcripts associated to one gene locus. Those transcripts came from several sources, namely mRNA sequences, but also from the EST database. Therefore, by searching UniGene database for UGT gene loci, we retrieved all associated transcripts, which have been then aligned between each others. By this simple exercise we sought to determine whether the UGT loci are associated with alternative mRNAs, i.e. transcripts having alternative exons1/promoters, internal exons or alternative ends.

The **Table 1** summarizes our findings, and presents new mRNAs for human *UGT* loci and supportive DNA clones. It is noteworthy that the transcript ID is an unofficial identifier only used to facilitate the description of each transcript. The latter includes sequences such as whole curated cDNA clones, but also single-passed sequences from either 5'-RACE, 3'-RACE, or shotgun-sequencing of cDNA clones. It is not surprising to observe alternative splicing events in all UGT loci, as presumed by the elevated ASE rate in human genes. The approach is solely qualitative and one may not deduce a relative abundance based upon the number of supportive sequences. We noted that ASEs mostly occur either at the beginning or the end of the mRNA molecules. Whether these mRNA species endogenously exist and generate proteins or, on the other hand, are modulator mRNA species (e.g. nonsense-mediated mRNA decay) have to be investigated. the deduced polypeptidic product for most ASEs would lead to shorter UGT proteins, each requiring molecular and biological characterization. One clear observation is the presence of diverse mRNA species likely having different functions and expression profile. Therefore, in future UGT expression experiment, researchers would face to clearly define the transcripts that would be addressed by their selected strategy to avoid further conflicting results. However, given the high sequence similarity between UGTs, this represents a significant challenge

Considering the already reported usage of alternative first exon in *UGT1A* locus, we only sought for ASEs in the common region. In fact, the use of an alternative exon 5 for UGT1A, previously described as exon 5b (Girard et al., 2007; Levesque, Girard,

Journault, Lepine, & Guillemette, 2007), is supported by five expressed sequences. It is noteworthy that one of these sequences is the reported mRNA from our previous published studies.(Girard et al., 2007; Levesque, Girard, Journault, Lepine, & Guillemette, 2007) Moreover, this new exon has been further investigated in more detail, as presented in a next section. No other ASE has been found. As abovementioned, in contrast to UGT1A locus, the UGT2 family members are encoded by single gene (Mackenzie, Bock et al., 2005; Riedy et al., 2000; Turgeon et al., 2000). We attempted to uncover ASEs associated to each gene. One major drawback of this approach is the high sequence similarity between either UGT2A or UGT2B. Indeed, the sequence quality of certain EST clones may impair the correct attribution to one gene cluster over another. In here, we aligned transcribed sequences associated with human UniGene clusters. This fast analysis allowed discovering several ASEs occurring in all UGT2B and UGT2A genes, excepted for UGT2B11, UGT2B15, and UGT2B17. We also have to note that splicing events occurring in the *UGT2B28* gene came from experimental evidence from the Dr. Alain Bélanger's laboratory (Levesque et al., 2001) and from several other monkey ESTs. The type of ASEs are various, we noted the use of either alternative 5' and 3' splice site (UGT2B4, UGT2B10, UGT2B28), the internal exon skipping (UGT2B4, UGT2B7, UGT2B10, UGT2B28), the alternative internal exon usage (UGT2B4). Although not alternative splicing events, we also noted the use of alternative promoter (alternative first exon in *UGT2B4* and *UGT2B7*), and alternative last exon (*UGT2B4* and UGT2B7). Some experimental evidences in support of these events have recently been obtained and are discussed below.

The major challenge is now to corroborate these new mRNA species in a cellular context, to define their abundance and expression profile, to determine their translational product (if it is so), and ultimately to establish their functional and physiological relevance. Firstly, assuming that these mRNAs may conduct to proteins, few of them should be functional transferase enzymes. These new translated polypeptides may serve for an unknown molecular process or act as a negative modulator of glucuronidation as discussed below. Otherwise, mRNA molecules bearing a prematured stop codon might intrinsically be involved in the nonsense-mediated mRNA decay (for review see (Chang, Imam, & Wilkinson, 2007). Several experimental studies are currently in progress and will be needed to clearly establish the role of all transcribed mRNA species, and whether these affect the net glucuronidation activity in a specific time and tissue.

5. Experimental evidence of new UGT1A-associated mRNAs

Extensive search for new putative exons at *UGT1* locus in several human tissues and cell lines was performed and led to discovery of a new exon, located in the common region of the gene, between coding exons 4 and 5 and was renamed exon 5b (**Figure 3**) (Levesque, Girard, Journault, Lepine, & Guillemette, 2007). This genomic sequence perfectly matches with the transcript UGT1A.2 reported in Table 1 found in EST database. However, according to our findings, this exon 5b might either be used as terminal exon (leading to the mRNA isoform variant 2 or v2) or be spliced with the common exon5 (exon5a) (leading to mRNA isoform variant 3 or v3)(**Figure 3**) (Girard et al., 2007). Thus, the discovery of the novel exon 5b initiated a search for a whole new class of UGT1A isoforms, and revealed the production of 18 new mRNAs (v2 and v3)

leading to 9 new UGT1A isoform 2 proteins (Girard et al., 2007; Levesque, Girard, Journault, Lepine, & Guillemette, 2007). The presence of exon 5b (v2 and v3) in the mRNA sequence generates a shorter polypeptide product (named isoform i2 or UGT1A i2) lacking the amino acids 435-533 encoded by exon 5a which are replaced by a sequence of 10 amino acids: RKKQQSGRQM (Girard et al., 2007; Levesque, Girard, Journault, Lepine, & Guillemette, 2007) (Figure 3). These new UGT splice variant isoforms 2 lack the transmembrane domain encoded by the exon 5a but possess an ERtargeting signal peptide and complete binding sites for both the substrate and cosubstrate. The heterologous expression of UGT1A i2 isoforms in HEK293 cells indicate that despite abundant expression and ER localization of i2 proteins, they are inactive in the transfer of the glucuronic acid moiety to classical substrates of the UGT1A family members. The observations that the splice i2 forms are devoid of transferase activity (Girard et al., 2007; Levesque, Girard, Journault, Lepine, & Guillemette, 2007) are in line with previous studies that reported the existence of a loss of enzyme activity for Cterminal truncated UGT proteins located in the ER (Levesque, Girard, Journault, Lepine, & Guillemette, 2007). However, they display a repressive effect on rates of formation of glucuronide products when co-expressed with their equivalent UGT1A il isoforms in HEK293 cells (Girard et al., 2007; Levesque, Girard, Journault, Lepine, & Guillemette, 2007). Indeed, co-expression of i1 and i2 within the cells result in significant reduction of glucuronidation activity of UGT1A1, UGT1A7, UGT1A8 and UGT1A9 (Girard et al., 2007). These data support a potential dominant negative role for these splice forms of the UGT1A gene.

Using available polyclonal anti-UGT antibodies, the presence of 45-kd UGT1A_i2 proteins was shown to be widely observed in human tissues, including liver, kidney, oesophagus, small intestine and colon (Girard et al., 2007). In most tissues, UGT1A_i2 are expressed together with their UGT1A_i1 homologs. From the published data, the expression of i2 was predominant in extrahepatic tissues compared to the liver. The subcellular localization of i2 to the endoplasmic and perinuclear structure was firstly described in UGT1A-overexpressing HEK293 cells together with classical isoforms i1 where protein interactions might occur (Levesque, Girard, Journault, Lepine, & Guillemette, 2007). The endogenous expression of splice i2 proteins in human-derived colon cell lines indicates its intracellular location to ER membrane and perinuclear structure, as demonstrated for the UGT1A isoforms 1. Immunohistochemal experiments in human colon tissues further support localization of both isoforms i1 and i2 in similar structures (Bellemare *et al.* under review).

Down-regulation of UGT1A_i2 proteins by transfecting specific small interference RNA into in human colon cells enhances the glucuronidation activity for various UGT1A substrates (Bellemare *et al.* under review). This supports the dominant negative hypothesis for UGT1A_i2 proteins. These results not only demonstrate the biological significance of alternative splicing of the UGT1A system, but also suggest a new UGT regulatory pathway. These observations clearly unveil another level of complexicity in the regulation of intracellular glucuronidation activity and may have important physiological and pharmacological implications. We propose that protein interaction may underlie the negative role of shorter i2 proteins, and there is substantial literature

supporting that UGT proteins might interact together as dimers/oligomers or with other ER-resident proteins (Finel & Kurkela, 2008; Fremont, Wang, & King, 2005; Fujiwara et al., 2009; Fujiwara, Nakajima, Yamanaka, Katoh, & Yokoi, 2007; Fujiwara et al., 2007; Ghosh et al., 2001; Ikushiro, Emi, & Iyanagi, 1997; Ishii et al., 2001; Kurkela et al., 2003; Kurkela, Hirvonen, Kostiainen, & Finel, 2004; Kurkela et al., 2007; Meech & Mackenzie, 1997b; Nakajima, Yamanaka, Fujiwara, Katoh, & Yokoi, 2007; Operana & Tukey, 2007; Radominska-Pandya, Ouzzine, Fournel-Gigleux, & Magdalou, 2005; Takeda et al., 2009). Interaction between both splice forms is also supported by their proximity in ER membranes, and by the demonstration of physical interaction between both isoforms 1 and 2 of UGT1A1 through co-immunoprecipitation experiments (Levesque, Girard, Journault, Lepine, & Guillemette, 2007). Together, these observations begin to highlight a dynamic and functional antagonism between isoforms 1 and their splice i2 variants. A parallel example was reported by Ghosh et al. in 2001, where an inactive truncated mutant was acting as a negative modulator of UGT1A1 activity while interacting with wild-type UGT1A1 (Ghosh et al., 2001). Further investigations are definitely required to provide molecular basis for these effects. Yet, such a negative regulation by differential oligomerization of the alternatively spliced isoforms is widely reported for a variety of proteins (Hashimoto, Chiorazzi, & Gregersen, 1995; Lincz et al., 2008; McElvaine & Mayo, 2006; Nakamura, Hashimoto, Furuyama, & Kakudo, 1995; Nakamura et al., 1995; Sztainberg et al., 2009; W. Wang, Kwok, & Chan, 2007).

The evolutionary conservation of a particular gene structure as well as of a specific splice isoforms is generally a good indicator of the functionality of those variants. Indeed, the structure of the *UGT1* gene is highly conserved among orthologues vertebrate's

genes; exon 5b is also present in primates and rodents leading to shorter splice variants corresponding to human i2 orthologous (Girard et al., 2007). Furthermore, the search in public databases established the conservation of isoform 2 splice variants (UGT1A5_v2: Genbank accession number XM001150762) in chimpanzee (*Pan troglodytes*). Thus, the alternative splicing of the last exon at *UGT1A* locus might be part of the tightly regulated circuit that controls the overall cellular glucuronidation activity.

These new alternative splicing events occurring in the common region of the UGT1A gene increase the protein diversity derived from this single gene, while representing new post-transcriptional and post-translational regulatory mechanisms of the glucuronidation pathway. Besides, it is well appreciated that alterations in splicing patterns of genes contribute to the regulation of gene function by generating endogenous inhibitor or activator molecules. This raises the possibility that changes in the equilibrium between endogenous i1 an i2 forms of UGT1A possibly represent a significant contribution in global conjugation activity of the cell. At present, the physiological and pathological roles of this particular class of *UGT1A* gene products remain an unanswered and open question. Preliminary results tend to show a strong heterogeneity in expression and localization of i2 within tissues, but several studies are needed to understand the exact cellular functions of these newly discovered splice variants. Understanding the regulation of this complex transcriptional system will be of great interest in the near future and may certainly underlie part of the interindividual glucuronidation variability currently unexplained by *UGT1A* single nucleotide polymorphisms or other non-genetic causes.

6. Alternative splicing at the *UGT2* gene loci.

6.1 The UGT2B28 gene

Although EST clustering analyses suggest alternative splicing events in human *UGT2* genes, there is still little experimental evidence of such events. Effectively, the scarce evidence of splicing in this subfamily comes initially from the description of the UGT2B28 gene in 2001 (Levesque et al., 2001). Indeed, two distinct alternatively spliced UGT2B28 transcripts were isolated and characterized at the molecular level along with the active UGT2B28 v1 transcript, originally described as UGT2B28 type I (Figure 4) (Levesque et al., 2001). The UGT2B28 type III transcript lacks a portion of the exon 1, potentially via the utilization of a cryptic splicing site, whereas the UGT2B28 type II lacks the exons 4 and 5 in the mature mRNA (Levesque et al., 2001). These transcripts were the first evidence for the existence of alternative splicing in this human UGT subfamily, and are related to the transcripts UGT2B28.2 and UGT2B28.3 in Table 1. Functional data performed with the corresponding cDNA revealed no evidence of transferase activity for the UGT2B28-derived protein variants when stably expressed independently in HEK293 cells. However, no co-expression experiments were performed in this previous study and the physiological relevance of these transcripts remained to be assessed.

6.2 The UGT2B7 gene

More recently, Innocenti and coworkers demonstrated the presence of two novel exons, called exon 1A and 1B in the *UGT2B7* gene upstream of the first coding exon (**Figure 4**)

(Innocenti et al., 2008). The authors showed the existence of two additional UGT2B7associated transcripts in human livers, of which one contain the exon 1A and 1B spliced with the coding exon 2 and 3 of UGT2B7 (UGT2B7 v2) and the other includes only the exon 1A joints to the coding exon 2 and 3 (UGT2B7 v3). They further reported that both isoforms ended in the intron 3. These isoforms are related to the predicted transcripts UGT2B7.3 and UGT2B7.4, respectively, described in Table 1, but, in contrast to Innocenti et al. (Innocenti et al., 2008), longer mRNA transcripts have been reported. Whether all these transcripts (shorter and longer) are produced by human cells remains to be uncovered. It is however clear that new UGT2B7 mRNAs are transcribed from an alternative promoter (upstream of the exon 1A). If these mRNAs were translated, it presumably start at the ATG in the coding exon 2, would lead to an UGT lacking the substrate-binding domain. Accordingly, whether the mRNA per se, the associated proteins, or both has functional relevance is still not known. Nevertheless, these observations will clearly influence the experimental design of subsequent studies about UGT2B7 gene expression and activity knowing that several mRNAs are produced by this very important metabolizing gene in the human liver and extrahepatic tissues.

6.3 The UGT2B4 gene: A closer look at the genomic level

To explore in more detail the formation of alternative splice products at the *UGT2* loci, and the validity of using a computational approach to predict mRNA transcripts from one locus, the *UGT2B4* gene was chosen as model (**Figure 4**). Based upon data in Table 1, we designed amplification strategies to both determine the presence of UGT2B4-associated ASEs and full-length isoforms. All predicted alternative transcripts and ASEs

listed in Table 1 were observed in human liver tissues. In addition, we uncovered other mRNA isoforms revealing that alternative splicing at the UGT2B4 locus produces significant gene expression diversity. Preliminary quantitative-PCR experiments show that splice variants are expressed at variable levels in the liver, whereas a more restricted pattern of expression seems to prevail in extrahepatic tissues (Lévesque, Ménard, Bellemare, Girard, & Guillemette). We further cloned and expressed individual splice UGT2B4 forms in HEK293 cells. Immunofluorescence analysis revealed that these transcripts conduct to proteins that are directed to the ER and perinuclear membranes despite lacking critical domains. Microsomal preparation from these stably transformed cells did not displayed detectable glucuronidation activity for eugenol, a broad substrate for UGT2Bs. However, in the presence of these new proteins, the rates of glucuronidation by the UGT2B4 i1 enzyme were significantly reduced by 30-50%. To further confirm the potential regulatory functions of these UGT2B4 splice proteins, additional experiments are required. However, as previously exemplified by the UGT1A v2/v3 alternative transcript, the production of various mRNA species by the UGT2B4 locus is not only related to erroneous transcription and/or splicing mechanisms, but is likely to have a functional role in the control of glucuronidation activity.

6.4 Conclusions on the topic of alternative splicing

Several questions remain to be answered before to substantiate functional role(s) for these new UGT transcripts *in vivo*. Nevertheless, we should now be aware of the presence of such transcripts, particularly in the design of future gene expression analysis. Among these questions are: what is their relative abundance in various tissues and conditions? Do their changes in expression significantly influence the glucuronidation? by which mechanism(s) and under which circumstances? Are splice variants associated with diseases or specific physiological conditions? Is *in vivo* synthesis of splice forms controlled, if so, by which mechanism(s)? Does splice variants contribute to variation in drug metabolism at the magnitude of clinically significance? Does this affect the way patients metabolize drugs or endogenous molecules? Several examples in the literature support that splice variants of genes may affect drugs metabolism and even drug resistance (reviewed by (Passetti, Ferreira, & Costa, 2009)). We might not promptly answer these questions, and several studies will be needed to understand the function and importance of UGT splice variants in cellular metabolism.

7. Copy-number variations of *UGT2B* genes

The human genome displays extensive copy-number variations (CNV) of large DNA fragments (McCarroll et al., 2006). In particular circumstance, this genomic instability might either induce gene duplication or deletion, and thus influencing the expression levels of the targeted gene product (Spielman et al., 2007). When associated to a *UGT* gene, it may impact upon interindividual drug metabolism variability. Indeed, one of the most studied CNV in human includes the sequence of the *UGT2B17* sex-steroid and drug metabolizing gene. This gene was isolated from the human prostate and the human prostate cancer cell lines in 1996 by Dr Belanger's group in the search of hormone-related cancer genes (Beaulieu, Levesque, Hum, & Belanger, 1996). UGT2B17 was then

shown to be involved in the termination of the androgen signal in target cells, conjugating and inactivating the major androgen dihydrotestosterone (DHT), testosterone and androsterone (Beaulieu, Levesque, Hum, & Belanger, 1996). Then, the discovery of a common deletion of the entirety of gene leading to CNV has been reported and is characterized by either the presence of one gene copy or the absence of the entire gene sequence (Jakobsson et al., 2006; Wilson et al., 2004). Subsequently, UGT2B17 has been showed to be one of the most commonly deleted genes in the human genome, this associated with the most significant variations in expression levels between ethnic groups (Jakobsson et al., 2006; McCarroll et al., 2006; Spielman et al., 2007). In addition, UGT2B17 was selected as an initial candidate to study the evolutionary divergence of copy-number variations (CNV) in human populations (Xue et al., 2008). The clinical interests for the deletion of UGT2B17 comes from its association with circulating testosterone levels (Jakobsson et al., 2006), its impact on doping in sports (Schulze et al., 2008), its association with prostate cancer risk revealed in some studies and its intrinsic role as an antigenic peptide in transplant procedures (Gallagher et al., 2008; Karypidis, Olsson, Andersson, Rane, & Ekstrom, 2008; Murata, Warren, & Riddell, 2003; Olsson et al., 2008; J. Park et al., 2006; J. Park et al., 2004; J. Y. Park et al., 2007; Xue et al., 2008). Moreover, CNV of UGT2B17 has recently been associated with the pathogenesis of osteoporosis and higher concentrations of circulating testosterone and estradiol and, therefore, represents a susceptibility gene for this highly prevalent metabolic bone disease in Asians (Yang et al., 2008). The UGT2B17-including CNV might in fact be seen as a human gene knockout enabling a more precise characterization of the physiological role of this gene in complex diseases including hormone-dependent cancers. Definitely, the

major advances in the understanding of the role of *UGT2B17* gene comes from CNV studies, which are critical for appreciating the significance of this particular gene product within this high homology subfamily of UGT2B enzymes.

To further emphasize the importance of CNV in this human subfamily, another CNV on chromosome 4q13 encompassing the *UGT2B28* gene has been found in human populations. *UGT2B28* is located 500 kb away from *UGT2B17* and is also involved in androgen and estrogen metabolism (Levesque et al., 2001). *UGT2B28* is the more recent characterized gene in this subfamily. As opposed to the well characterized impact of *UGT2B17* deletion in human populations, the role of *UGT2B28* CNVs has never been addressed. Nevertheless, a recent association study using tag-SNPs (associated with the *UGT2B28* genomic deletion) suggested that foetal development and gestation length are influenced by the presence of *UGT2B28* CNVs; a feature which may be related to its detoxification function or its role in androgen and/or estrogen metabolism (C. L. Relton, 2008). In support, changes in expression levels of *UGT2B28* was also associated with progression of oesophageal dysplasia in a Chinese chemoprevention trial (Joshi et al., 2006), underscoring the needs to study *UGT2B28* CNVs as well.

To facilitate genomic studies in this high homology region on chromosome 4q13, the molecular identification of the breakpoints of these two common deletions was localized precisely (Menard, Eap, Harvey, Guillemette, & Lévesque, In press). Segmental duplications of 4.9 kb for *UGT2B17* and 6.8 kb for *UGT2B28* are localized 117 kb and 108 kb apart respectively on both ends of the deletions and both comprise purine-rich

recombination sites. CNVs of UGT2B17 and UGT2B28 occur at a frequency in Caucasians of 27% and 13.5%, respectively. We also demonstrate that only 43% of individuals have 2 copies of both genes, whereas 57% harbour at least one deletion. Their co-occurrence on 5% of the chromosomes creates a large genomic gap of 225 kb in these individuals. Their co-existence with a functional polymorphism of the androgenmetabolizing gene *UGT2B15*, p.D85Y (rs1902023:G>T) in Caucasians is further shown. Haplotype analysis revealed that the long arm of chromosome 4 can harbour seven different haplotypes, the 2B17+/2B28+/2B15Y⁸⁵ (Hap I), 2B17+/2B28+/2B15D⁸⁵ (Hap II), 2B17-/2B28+/2B15D⁸⁵ (Hap III), 2B17+/2B28-/2B15Y⁸⁵ (Hap IV), /2B28+/2B15Y⁸⁵ (Hap V), 2B17-/2B28-/2B15D/Y⁸⁵ (Hap VI and VII) with frequencies of 34.5%, 29.5%, 15.5% and 8.5%, 7% and 5% respectively (Figure 5) (Menard, Eap, Harvey, Guillemette, & Lévesque, In press). The exact haplotypes frequencies of these three metabolizing genes in other ethnic group remains undetermined but is expected to be different because of the greater prevalence of UGT2B CNVs in other populations compared to Caucasians (McCarroll et al., 2006). The implications of these findings are critical because the polymorphic deletions of UGT2B17 and UGT2B28 are associated with one of the UGT2B15 codon 85 allele, which can have a profound impact on steroids metabolism in presence of CNVs in both these genes. Of note, in the last years, several studies were performed with respect to the UGT2B17 deletion genotypes mainly in prostate cancer patients with conflicting results (Gallagher et al., 2008; Karypidis, Olsson, Andersson, Rane, & Ekstrom, 2008; J. Park et al., 2006; J. Park et al., 2004; J. Y. Park et al., 2007). CNVs of UGT2B17 and UGT2B28, their distinct associations with a polymorphism of the UGT2B15 gene and their variable prevalence in diverse ethnic

groups may indeed lead to divergent conclusions. Furthermore, several studies demonstrate the impact of CNV of *UGT2B17* on hormonal status in humans; however, the co-existence of *UGT2B17* and *UGT2B28* CNVs in 15% of the studied cohort is likely to influence the overall sex steroid metabolism in these subjects, the latter genes being involved in both androgens and estrogens elimination. Therefore, the genomic status of all these steroid metabolizing genes should be evaluated systematically in order to refine the molecular understanding of this complex locus and their physiological consequences on the endocrine system. Based on our data, it seems clear that haplotype analyses are critical because co-existence of these two common major deletion polymorphisms, in conjunction with their genetic associations with SNPs in the *UGT2B15* gene, may have even more profound physiological consequences on the overall hormonal metabolism then each of these functional polymorphisms taken individually.

8. Epigenetic regulation of *UGT* gene expression.

Epigenetic regulation such as DNA methylation is another key mechanism to either activate or silence gene transcription, and abnormal epigenetic regulation has been described as an important characteristic of tumor progression and malignancy (Grady & Carethers, 2008; Shelton, Misso, Shaw, Arthaningtyas, & Bhoola, 2008). Epigenetic mechanisms are also important to maintain tissue-specific gene expression. Furthermore, abnormal methylation of genes is by far more frequent and has often more drastic consequences than classical genetic mutation (Al-Romaih et al., 2007). Currently few evidence support the epigenetic modulation of human UGT expression and one of the few published example is the significant interindividual variability in the level of

methylation of the *UGT1A1* gene in cellular models derived from human colon tumors. UGT1A1 is well known to be responsible for the inactivation of SN-38 (Gagne et al., 2002), the active metabolite of the prodrug irinotecan use to treat metastatic colorectal cancer. Therefore, any epigenetic mechanisms altering the UGT1A1 gene has the potential to modify the glucuronidation capacity at the target cell level and, hence, the biological effect of the drug. Aberrant methylation of specific CpG islands of the UGT1A1 gene in UGT1A1-negative cells (HCT116, HCT-15, and COLO-320DM). Data support that UGT1A1 promoter hypermethylation results in complete repression of gene transcriptional activity (Gagnon, Bernard, Villeneuve, Tetu, & Guillemette, 2006). Reversal of promoter methylation by treatment with demethylating and histone deacetylase inhibitor agents had the capacity to restore UGT1A1 expression in hypermethylated UGT1A1-negative cells. Loss of UGT1A1 methylation was further associated with an increase in UGT1A1 protein content and with an enhanced inactivation of SN-38 by 300% in HCT116 colon cancer cells (Gagnon, Bernard, Villeneuve, Tetu, & Guillemette, 2006). An analysis of the *UGT1A1* 5'-flanking sequence (−540 to −1) in human colon cancer cells has revealed that hypomethylation of this region is important for UGT1A1 transcription (Gagnon, Bernard, Villeneuve, Tetu, & Guillemette, 2006). More specifically, the extent of *UGT1A1* promoter methylation between CpG -1 (-4nt relative to the ATG) and -4 (-99nt relative to the ATG) of the promoter was shown to significantly predict *UGT1A1* gene expression in colon cancer cell lines (Gagnon, Bernard, Villeneuve, Tetu, & Guillemette, 2006). Based on the above findings, it seems that epigenetic modifications are part of the complexity underlying UGT gene expression. At this point, it remains an open question whether the expression of additional UGTs is regulated by epigenetic mechanism in a tissue-specific manner and if such regulation has phenotypic consequences.

9. Concluding Remarks

Over the last few years, the emphasis was directed towards the characterization of genetic variations in all previously characterized *UGT* genes in the hope of understanding the considerable genetic variability observed for this major molecular pathway. Despite finding some hints, the global genetic plasticity of the glucuronidation pathway remains unsolved. As discussed here, part of this diversity is expected to occur via other genomic processes, including the production of alternative transcripts, copy-number variations and epigenetic processes (Figure 6). Alternative splicing occurs in most genes and the UGT genes are definitely not an exception to this principle. At present, despite the fact that very few studies have been conducted, it is evident that extensive splicing patterns occur in the *UGT* genes and these processes are just beginning to emerge especially for *UGT*2 family members. Predicted transcripts from public databases and experimental analyses undoubtedly sustain the presence of alternative transcripts in *UGT* genes, and at present, this should change the design of future experiment particularly regarding gene expression. Altogether, these cellular processes drastically increase the genetic variability arising from the UGT1A and UGT2B genes and protein diversity. By these observations, the analysis of high interindividual variability and diversity of the glucuronidation pathway is becoming even more complex and several years of investigation will certainly be needed to understand the physiological and pharmacological roles of these new processes.

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Figure Legends

Figure 1. Schematic representation of the mRNA diversity generate by known alternative splicing patterns. Some of these splicing events have been observed in UGTs.

Figure 2. A) Phylogram of the human UGT1A and UGT2B proteins and their corresponding cluster of genes on chromosomes 2 and 4, respectively. Schematic representation of the UGT1A9 (B) and UGT2B7 (C) protein primary structures derived from each locus. Similarities between the two classes of proteins are shown.

Figure 3. Schematic representation of the *UGT1A* gene and the alternative sharing and splicing events taking place at the 5' and 3' ends of the locus. (Top) The previous genomic structure of *UGT1A* included 17 exons: 13 first exons and four common exons (2–5a). We found that the *UGT1A* locus encodes one additional exon (referred to as the new exon 5b) and that the common region comprises five shared exons instead of four, as currently reported in the literature. (Bottom) The newly identified *UGT1A* gene products (mRNAs) are referred to as variants, or v, whereas new proteins are named isoforms, or UGT1A_i, according to Human Gene Nomenclature Guidelines. The previously described 'classical' UGT1As are named UGT1A_i1 and the newly described UGT1A proteins are referred to as UGT1A_i2. Exon 5a and the new exon 5b are alternatively spliced, generating variant UGT1A mRNAs: v1 (containing 5a), v2 (5b) and v3 (5a+5b). The proteins translated from v3 are identical to those encoded by the UGT1A_v2 mRNAs, as they have identical open reading frames. These data suggest the existence of nine additional UGT1A_i2 proteins, increasing to 18 the number of proteins encoded by UGT1.

Figure 4. Known Alternatively spliced UGT2B mRNA transcripts: Depicted here are distinct mRNA transcripts which were discovered by our group and/or be found on the NCBI website. The mRNA sequences of these transcripts are present in the Genbank database, the identification number on the website is also indicated.

Figure 5. Haplotypes of the *UGT2B17*, *UGT2B28* CNV along with the *UGT2B15* polymorphism on chromosome 4. A total of seven different haplotypes of *UGT2B17*, *UGT2B28* and *UGT2B15* have been inferred, using genotyping data. White boxes represent wild-type state (no polymorphism), while deletions are depicted by a crossed-out shaded box and variant nucleotides by a red box.

Figure 6. Current and emerging genomic mechanisms likely underlying interindividual glucuronidation phenotypes.

Table 1. Predicted transcripts from EST clusters at the UGT1 and UGT2 loci.

Locus (Unigene cluster) (coding strand) ¹	Number of Associated transcripts ²	Transcript ID ³	Exon Boundaries ⁴	Supportive Seq ⁵ (gi number)	Comments
UGT1A (Hs.554822) (+ strand)	404	UGT1A.1	182291-182422 183106-183193 183477-183696 187579-187816		Only Common exon 2 to 5
		UGT1A.2	182291-182422 183106-183193 183477-183696 184788-185275	19140059 19010676 12671933 31657195	Use of an alternative last exon (exon 5b).
UGT2B4 (Hs.285887) (- strand)	78	UGT2B4.1	1020761-1020006 1018707-1018559 1014436-1014305 1011562-1011475 1010293-1010074 1005776-1005031		Total protein length= 529 aa.
		UGT2B4.2	1050880-1050553 1020276-1020006 1018707-1018559 1014436-1014305 1011562-1011475 1010293-1010074 1005776-1005031	194390249 146076957	A novel first exon is spliced at an alternative 3' splice acceptor site in coding exon 1. Deletion of aa 1 to 150. New ATG start codon in the novel first exon encoding for a new peptide MFFALLHVSSTNGL in frame with the remaining UGT2B4 protein (aa 151-529).
		UGT2B4.3	1020761-1020006 1018707-1018559 1014436-1014305	37777528	The insertion of an alternative exon between coding exon5 and 6. Prematured ending of the proteins

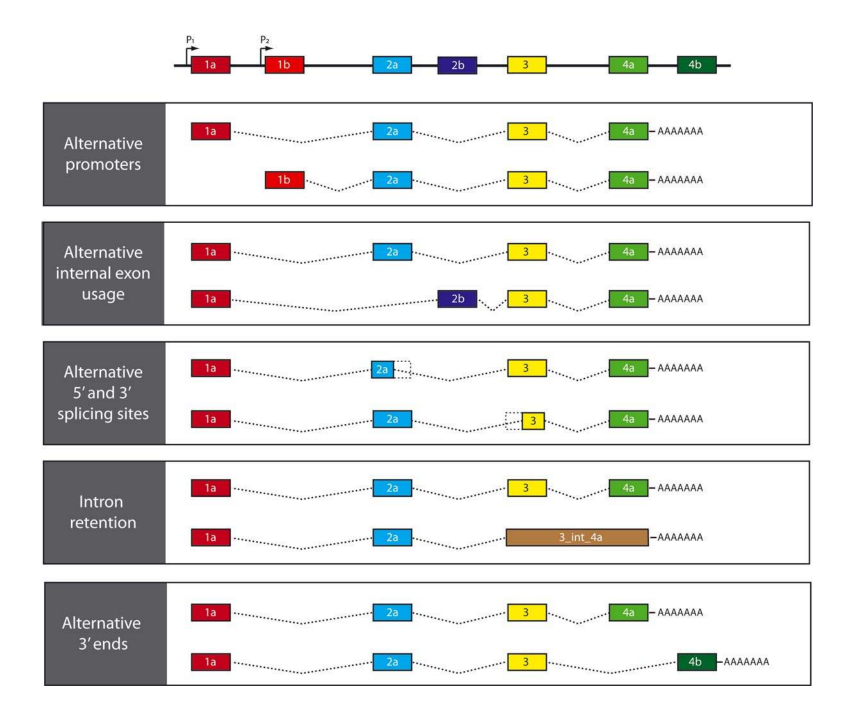
			1011562-1011475 1010293-1010074 1006131-1005990 1005776-1005031		at aa 436.
		UGT2B4.4	1020761- 1020259 1018707-1018559 1014436-1014305 1011562-1011475 1005776-1005031	31023872 37777529	Alternative 5' donor splice site in the exon1, and skipping of the coding exon5. Prematured ending of the proteins at aa 157 followed by 82 UGT2B4-unrelated amino acids.
		UGT2B4_ASE.1	1050880-1050553 1020832-1020006	80808841	Alternative 5'UT-exon, which is spliced with the first coding exon. Full-length UGT2B4 protein is expected.
		UGT2B4_ASE.2	1050880-1050553 1018707-1018559	52172943	Alternative 5'UT-exon which is spliced with the coding exon2. The translation is presumed to start in the novel first exon.
UGT2B7 (Hs.654424) (+ strand)	251	UGT2B7.1	621373-622107 623406-623554 627673-627804 632041-632128 632969-633188 637323-637853		Total protein length= 529 aa.
		UGT2B7.2	621373-622107 623406-623554 627673-627804 632041-632128 637323-637853	10735032	Deletion of the coding exon5. Prematured ending of the proteins at aa 363.
		UGT2B7.3	575965-576468 614338-614469	13333538 28109514	Two alternative 5'UT-exons which are spliced with the coding exon2.

			623406-623554 627673-627804 632041-632128 632969-633188 637323-637853	28111933 15748889 13582705	The translation is presumed to start in the coding exon 2, and would lead to UGT2B7 lacking the substrate binding domain.
		UGT2B7.4	575965-576468 623406-623554 627673-627804 632041-632128 632969-633188 637323-637853	12673874	One alternative 5'UT-exon spliced with the coding exon2. The translation is presumed to start in the coding exon 2 as abovementioned.
UGT2B10 (Hs.201634) (+ strand)	75	UGT2B10.1	340861-341603 342895-343042 347136-347268 351276-351363 352195-352414 355466-355745		Total protein length=529 aa.
		UGT2B10.2	340861- 341351 342895-343042 347136-347268 351276-351363 352195-352414 355466-355745	22119058 37777525 194391069	Alternative 5' donor splice site in the coding exon1. Do not change the coding frame. Deletion of aa 156 to 239.
		UGT2B10_ASE.1	340861- 341038 342895-343042	52111972	Alternative 5' donor splice site in the coding exon1. Change in the coding frame and Prematured ending of the proteins.
		UGT2B10_ASE.2	351276-351363	13715524	Deletion of the coding exon5.

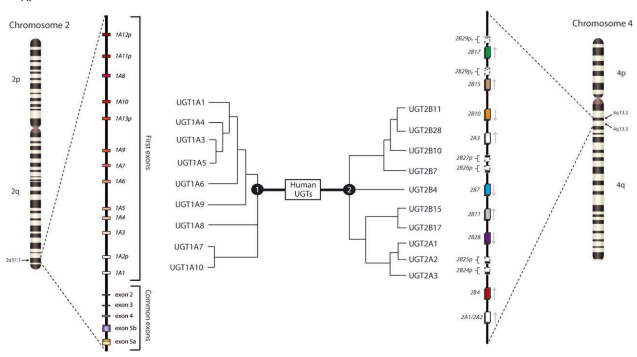
			355466-355745		
UGT2B28 (Hs.653154) (+ strand)	9	UGT2B28.1	805365-806087 807380-807528 811618-811749 814531-814618 815458-815677 819396-819675		Total protein length= 529 aa.
		UGT2B28.2	805365-806087 (□ 805685-806038) 807380-807528 811618-811749 814531-814618 815458-815677 819396-819675		Deletion of aa 107-224 in the coding exon1.
		UGT2B28.3	805365-806087 807380-807528 811618-811749 819396-819675		Deletion of the coding exon4 and 5.
UGT2A1(2A2) (Hs.225950) (- strand)	21	UGT2A1.1	1172566-1171796 1124260-1124112 1121247-1121116 1120115-1120028 1119562-1119343 1114517-1113283		Total protein length= 527 aa.
		UGT2A1.2	1172566-1171796 1160688-1160557 1124260-1124112 1120115-1120028 1119562-1119343	148753379	The use of an alternative exon between the coding exon1 and 2. Deletion of the coding exon3.

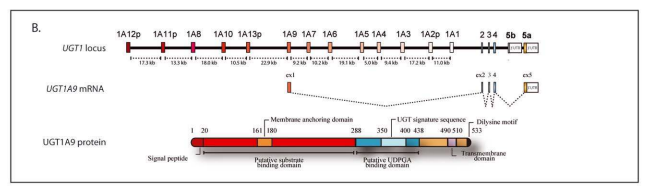
		UGT2A2.1	1114517-1113283 1164482-1163765 1124260-1124112 1121247-1121116 1120115-1120028 1119562-1119343 1114517-1113283		Total protein length= 528 aa.
UGT2A3 (Hs.122583) (- strand)	64	UGT2A3.1 UGT2A3_ASE.1	476698-475912 470320-470172 457625-457494 456108-456021 455631-455412 454958-453326 470320-470172 468244-468148 457625-457494 456108-456021 455631-455412	194373594	Total protein length= 527 aa. Alternative exon between exon 2 and 3.

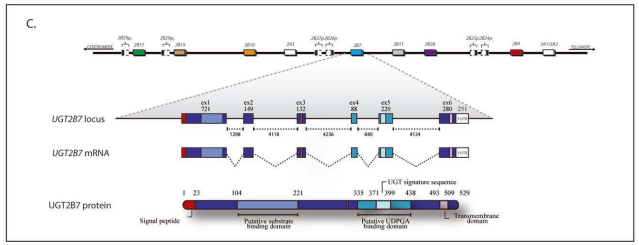
- 1. The coding strand indicates the orientation of the transcription, (+) from centromere to telomere, (-) from telomere to centromere.
- 2. The number of transcripts is the total number of mRNA and EST sequences associated with the corresponding locus in the Unigene cluster.
- 3. It is noteworthy that the transcript ID is an unofficial identifier only used to facilitate the description of each transcript. The transcript ID UGTXXX.X, indicate a full-length mRNA. The transcript ID UGTXXX_ASE.X indicate only the observation of an alternative splicing event (ASE).
- 4. Nucleotide position is given according to the human genome contig NG_002601 for UGT1A locus, and the contig NT_077444 for all UGT2 genes.
- 5. The supportive sequences for the major mRNA transcript for each locus, UGTXXX.1, have not been listed.



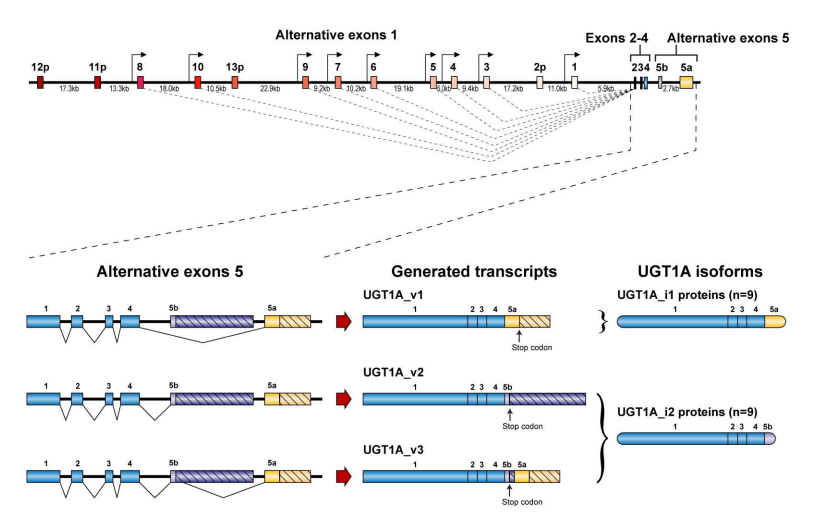


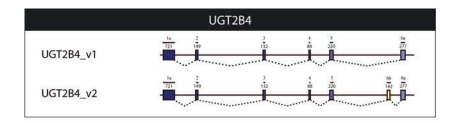


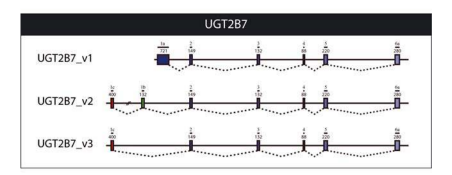


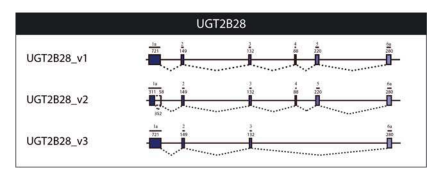


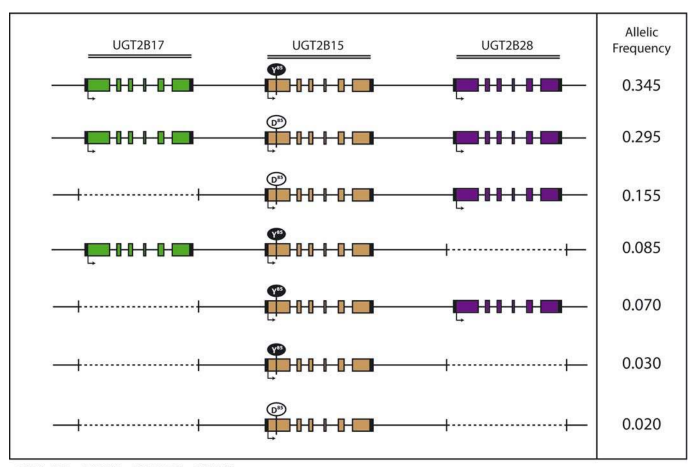
UGT1A Locus











 $2B15_D^{85} = 2B15*1, \ 2B15_Y^{85} = 2B15*2$