Identification of N-Acetyltransferase 2 (NAT2) Transcription Start Sites and
Quantitation of NAT2-specific mRNA in Human Tissues

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ABBREVIATIONS: NAT2; N-acetyltransferase 2; NAT1, N-acetyltransferase 1; RT, reverse transcriptase; PCR, polymerase chain reaction; q-RT-PCR, quantitative reverse transcriptase polymerase chain reaction; ORF, open reading frame; 5’-RLM-RACE, 5’-RNA ligase-mediated rapid amplification of cDNA ends; TSS, transcription start sites; BD, Biosciences Clontech.
ABSTRACT

Human N-acetyltransferase 2 (NAT2) genetic polymorphism is associated with drug toxicity and/or carcinogenesis in various tissues. Knowledge of NAT2 gene structure and expression are critical for understanding these associations. Previous findings suggest that human NAT2 expression is highest in liver and gut, but expressed at functional levels in other tissues. A sensitive and specific TaqMan reverse transcriptase polymerase chain reaction (RT-PCR) assay with intron-spanning primers was developed and used, together with a second TaqMan RT-PCR assay based on amplification of a NAT2 open reading frame (ORF) exon segment, to measure NAT2 mRNA in 29 different human tissues. Cap-dependent amplification of mRNA 5’ termini and review of public database information was done to more precisely define the NAT2 promoter(s) and to validate the quantitative RT-PCR assay design. The great majority (40/41) of NAT2 liver cDNAs had 5’ termini between 8682 and 8752 nucleotides upstream of the NAT2 ORF exon, and 34/40 5’-termini were at the -8711 and -8716 adenines. All of 59 NAT2 cDNAs with 5’ termini in this vicinity, including 40 of the liver isolates and 19 cDNAs in public databases from liver and other sources, showed direct splicing to the ORF exon, with no other non-coding exon detected. NAT2 mRNA was highest in liver, small intestine and colon and readily detected in most other tissues albeit at much lower levels. NAT2 expression in diverse human tissues provides further mechanistic support underlying associations between NAT2 genetic polymorphism, drug toxicity and/or chemical carcinogenesis.
Genetic polymorphism of the N-acetyltransferase 2 gene (NAT2) is strongly implicated in differential susceptibility to adverse drug reactions (Weber and Hein, 1985; Butcher et al., 2002) and to various diseases (Boukouvala and Fakis, 2005) especially cancers of the urinary bladder (Garcia-Closas et al., 2005; Carreon et al., 2006; Hein, 2006) and colon (Lilla et al., 2006; Moslehi et al., 2006) upon exposure to carbo- and heterocyclic-aromatic amine carcinogens. A large number of studies have also examined the possible involvement of NAT2 in the etiology of cancers of various other organs (reviewed in Hein et al., 2000a,b). A critical question is whether vulnerability to neoplastic transformation is related to specific expression of NAT2 in the target organ (Williams, 2001) and studies have investigated NAT2 expression in various human (reviewed in Boukouvala and Fakis, 2005) and recently in Syrian hamster (Hein et al., 2006) and mouse (Loehle et al., 2006) tissues. We hypothesized that human NAT2 expression is highest in liver and gut, but is also present in other human tissues. To test this hypothesis, sensitive, specific and well-characterized quantitative mRNA assays were used to quantitate relative expression of NAT2 in 29 different human tissue types.

Accurate quantitative measurements of NAT2 mRNA and protein pose a particular technical challenge because of the presence of the paralogous gene, NAT1, which is ubiquitously expressed in human tissues and shares 87% nucleic acid homology within the protein coding region and 80% amino acid identity to NAT2 (Blum et al., 1990; Ohsako and Deguchi, 1990). The high degree of nucleic acid homology between NAT1 and NAT2 is a potential confounder of general microarray-based analyses of mRNA in
normal human tissues. Thus, the reliability of NAT2 and NAT1 expression data in public databases such as GEO Profiles (http://www.ncbi.nlm.nih.gov/) or from genome-wide scans (Shyamsundar et al., 2005) is dependent on factors which may not always be consistent, including the homology of the chosen probe to both NAT1 and NAT2 mRNAs and the experimental hybridization stringency. Although the NAT1 and NAT2 enzymes are distinguishable by differences in substrate selectivity and affinity, the utility of activity measurements for quantitative evaluation of multiple human tissues is constrained by assay sensitivity as well as the requirement for fresh tissue specimens with functionally intact enzyme from individuals with the same polymorphic genotype. The use of mRNA detection methods with validated specificity thus offers the most practical and reliable means for sensitive quantitation of the relative expression levels of NAT2 in diverse human tissues.

Recent reports have shown that NAT1 gene transcription involves at least two different promoters and more than 10 different 5’ non-coding exons (Husain et al., 2004; Boukouvala and Sim, 2005; Butcher et al., 2005; Barker et al., 2006). Two previous studies indicated that NAT2 mRNA synthesis is less complex, involving transcription from a single promoter located more than 8.7 kb upstream of the ORF-exon and removal of a single 8.7 kb intron during splicing (Ebisawa and Deguchi, 1991; Deguchi, 1992), but tentative evidence for an additional promoter site was recently described (Boukouvala and Sim, 2005). To critically test the major single-promoter model of NAT2, we have now conducted an extensive 5’-RLM-RACE analysis of liver RNA and compared the
results of quantitative reverse transcriptase PCR (q-RT-PCR) assays based on intron-spanning versus intra-exon primer sets in diverse tissues.
MATERIALS AND METHODS

5'-RLM-RACE analysis of NAT2 mRNA. 5'-RLM-RACE analysis was performed starting with 10 µg total RNA, using the First-Choice RLM-RACE kit (Ambion) according to the manufacturer’s instructions. The final reverse transcription (RT) was performed with random decamer primers in a 20 µl reaction. The primers used for nested PCR were the adapter primers supplied by Ambion and the NAT2-specific outer, 5’-TTCCTCTCTCTTTCTGTCAAGCAGAAAATG-3’, and inner, 5’-AACCATGCCAGTGCTGTATTTGTAA-3’, primers. The NAT2 primers were designed to be specific for NAT2 (not NAT1) and to avoid the positions of known polymorphic sites. To obtain clones representing independently initiated transcripts, multiple independent nested PCRs were carried out using as template 0.5 or 1 µl of product from the RT reaction stage of the 5'-RLM-RACE procedure with liver RNA. The outer reactions were each 25 µl. Two µl of the resulting products were templates for subsequent 30 µl inner reactions. All PCRs were started in a preheated 94°C block followed by 94°C for 3 min, and 35 cycles of 94°C for 30 s, 60°C for 30 s, 72°C for 90 s with a final extension at 72°C for 7 min. PCR products for sequencing were resolved on a 1% agarose gel, purified with Qiaex II (Qiagen) and cloned with the TOPO TA Cloning Kit into the pcDNA 3.1/V5-His-TOPO vector (Invitrogen). Sequence reactions prepared with ABI Big Dye Terminators were analyzed on the ABI 310 Genetic Analyzer, as recommended by the manufacturer (Applied Biosystems). Transcription start sites were inferred from sequence adjoining that of the 5'-RLM-RACE adapter and mRNA splice locations were identified by alignment of cDNA sequences with NAT2 genomic
sequences using the BLAT tool (Kent, 2002) at the UCSC Gateway (http://genome.ucsc.edu/cgi-bin/hgGateway).

**Database analyses.** NAT2-specific cDNA and spliced EST sequences were identified by searches at the NCBI Unigene (http://www.ncbi.nlm.nih.gov/UniGene), DBTSS (http://dbtss.hgc.jp) and UCSC Gateway databases (Kent et al., 2002; Suzuki et al., 2002; Wheeler et al., 2006; Yamashita et al., 2006). Selected non-redundant cDNA sequences were aligned with genomic DNA using the BLAT tool (Kent, 2002) at the UCSC Gateway.

**Human RNA samples.** Purified total RNAs from human tissues were obtained from two independent commercial sources, Ambion (Ambion Inc., Austin, TX) and BD (BD Biosciences Clontech, Palo Alto, CA). The human tissue expression panel from Ambion included adipose, bladder, brain, cervix, colon, esophagus, heart, kidney, liver, lung, ovary, placenta, prostate, skeletal muscle, small intestine, spleen, testis, thymus, thyroid and trachea. Each of the Ambion panel RNAs was a pool derived from three individuals. A sample of breast RNA, also obtained from Ambion, was derived from one female. The expression panel obtained from BD included RNAs from brain, cerebellum, colon, fetal brain, fetal liver, heart, kidney, lung, placenta, prostate, salivary gland, skeletal muscle, small intestine, spinal cord, spleen, stomach, testis, thymus, trachea and uterus. The BD panel RNAs generally were pooled from a variable number of individuals, ranging from 3 to 63, however the brain, heart, placenta, and stomach RNAs were all derived from single individuals. RNAs representing adrenal gland, bladder and liver were also obtained from BD. For the 5’-RLM-RACE analysis, the liver RNA sample from BD and a second liver
RNA sample from Ambion, representing different individual males were used. MCF-7 cells were grown as described previously (Husain et al. 2004). HepG2 human hepatocellular carcinoma cells were obtained from the ATCC(# HB-8065) and cultured in DMEM/F12 (1:1) (Gibco) supplemented with 10% fetal bovine serum (Hyclone), penicillin (100 U/ml) and streptomycin (100 ug/ml) (BioWhittaker). Cells were propagated in a 37°C humidified chamber containing 5% CO₂. Cell culture medium was routinely changed every 2-3 days. Frozen samples of human colon tumor cell lines Caco-2 (J.C. States), SW480 (W.S. Zundel) and HT-29 (D.M. Miller), human lung adenocarcinoma cell line A549 (C.M. Klinge), and human embryonic kidney-derived cell line HEK293 (C.M. Klinge) were provided by the laboratories of the indicated University of Louisville investigators. RNAs were extracted as previously described (Barker et al., 2006).

**Quantitative real-time RT-PCR.** For quantitative real-time RT-PCR, TaqMan analysis was performed using the 7700 sequence detection system from ABI (Applied Biosystems, Foster City, CA). The 20 µl PCR reactions were 1X TaqMan Universal Master Mix, with *NAT2* forward and reverse primers at 300 nM and 100 nM probe. The primers, 5’-TTGGAAGCAAGAGGATTGCAT-3’ and, 5’-GATCTGGTGCTCAAGAATGTCAGT-3’ span the position of the 8.7 kb intron (Ebisawa and Deguchi, 1991; Deguchi, 1992; Boukouvala and Sim, 2005) and were used with the probe 6FAM 5’-TCCTGGTTGCTGGCC-3’MGBNFQ. A second *NAT2* primer pair and probe set has been described (Zang et al., 2004) and is located entirely within the 3’ portion of the single exon that contains the entire *NAT2* ORF. For PCR, initial
incubations at 50°C for 2 min and 94°C for 10 min were followed by 40 cycles of 95°C for 15 s and 60°C for 1 min. Quantitation of the endogenous control 18S rRNA was performed using TaqMan Ribosomal RNA Control Reagents for 18S rRNA (Applied Biosystems). Four µl of diluted cDNA, equivalent to 40 ng of the initial RNA template, was used in each PCR. Two independent RT reactions were prepared for each RNA, and for each cDNA product, real-time PCRs were carried out in duplicate, for a total of four readings. Each assay plate included reaction mix controls with water in place of template. As an additional negative control, TaqMan quantitation of reactions prepared in parallel without addition of RT were also run at least once for each set of replicate RT reactions. Baselines and threshold levels were selected as recommended by ABI and values of Ct, the cycle number at which the measured fluorescence reached the set threshold value, were recorded. For each real-time run, a ∆Ct (\(NAT2\) Ct – 18S rRNA Ct) was calculated for each sample and a ∆∆Ct value was then derived by subtracting the smallest ∆Ct from all ∆Ct values. The formula \(2^{(-\Delta\Delta Ct)}\) was used to calculate an initial relative concentration value for all samples, except that samples with a \(NAT2\) Ct of 40 were assigned as zero. For each RT set, the measures obtained from the two real-time runs were averaged and normalized with respect to the average of all samples in the same RT group.
RESULTS

Molecular analysis of the 5' structure of NAT2 mRNA. Two human male liver RNA samples, obtained from independent commercial sources, were each submitted to 5'-RLM-RACE for identification of NAT2 transcription start sites (TSSs) and determination of 5' mRNA splicing patterns. Clones of 41 independent 5'-RLM-RACE products were sequenced. The location of 40 of the TSSs was between 8,682 and 8753 bp upstream of the NAT2 ORF exon (Figure 1A), with 9 TSSs at the -8711 adenine and 25 TSSs at the -8716 adenine. These two adenines were located at positions -59 and -64 with respect to the splice donor sequence utilized in all of the corresponding mRNAs. The only atypical 5'-RLM-RACE clone originated from a start site located a further 4.6 kb upstream from the -8.7 kb promoter and the spliced structure did not include any sequences from the -8.7 kb promoter region (data not shown).

Figure 1B summarizes the positions of the 5' nucleotides present in 19 different NAT2 cDNA clone sequences retrieved from the public databases described in Materials and Methods. Although the cloning of most of these cDNAs was not specifically intended to detect TSSs, every observed 5' terminus lies within the same ~70 bp region as the TSSs identified by 5'-RLM-RACE, and a similar pattern of clustering of 5' ends in or near the -59 to -64 segment is apparent. The three clones from DBTSS, HSI05750, HSI08034 and HSI15615, were isolated from small intestine RNA with an “oligo-capping” method (Suzuki et al., 2002) which, like 5'-RLM-RACE, defines the position of capped 5' nucleotides. The 5' termini of two of the small intestine DBTSS clones are at -64, which is also the most common TSS in liver, and the third is at -61.
The region containing the \textit{NAT2} TSSs has 85\% sequence identity to the major transcription start site region of the paralogous \textit{NAT1} gene (Figure 1C). Although there was more diversity in the locations of \textit{NAT1} TSSs identified in breast tissue and a breast tumor cell line (Husain et al., 2004), one of the frequent \textit{NAT1} TSSs occurred at an adenine which corresponds to the primary \textit{NAT2} TSS adenine at the -64 position (Figure 1).

For all 40 of the liver 5'-RLM-RACE cDNAs with 5' termini in the -8.7 kb promoter region, the mRNA splicing pattern conformed to that shown in Figure 2, utilizing the consensus splice donor sequence located 8,658 nucleotides upstream of the \textit{NAT2} ORF exon (Figure 1). All 19 of the \textit{NAT2} public database cDNAs also utilize this splice donor sequence, and 18/19 are spliced to the typical \textit{NAT2} ORF exon acceptor site. One cDNA, BG569272, exhibits splicing which eliminates 339 nucleotides of the \textit{NAT2} ORF exon and may represent an aberrant transcript form or cloning artifact.

\textbf{Quantitative RT-PCR analysis of \textit{NAT2} mRNA in diverse human tissues.} Two different TaqMan primer and probe sets were utilized for quantitative analysis of \textit{NAT2} mRNA (Figure 2). In genomic DNA, the primers of the intron spanning assay set, IS-for and IS-rev (Figure 2), are separated by the 8.7 kb intron, eliminating any measurement error due to possible DNA impurities in RNA samples. The IS-for primer sequence is entirely 3’ to 38/40 of the \textit{NAT2} TSS locations identified in liver RNA (Figure 1A) and all 3 of the authentic TSSs from small intestine (Figure 1B). TaqMan intron-spanning assay results, on a logarithmic scale, are shown in Figures 3A and 4A for two sets of
human RNA samples representing 29 different tissue types, including 15 tissues represented in both panels. In each set, the highest NAT2 mRNA level was found in liver, with similar or slightly less expression found in small intestine and about 5-10 fold lower expression in colon. For several tissues, including kidney, lung, spleen and testis, which are represented in both the Ambion (Figure 3A) and the BD series (Figure 4A), NAT2 expression levels were consistently about 1% of that detected in liver. Lower or more variable levels of expression occurred in several other tissues, and consistently low levels of NAT2 expression were found in both of the samples representing heart, skeletal muscle, thymus and trachea (Figures 3A and 4A).

Figures 3B and 4B show results of a second NAT2-specific TaqMan q-RT-PCR assay, which utilizes primers and a probe all located within the single NAT2 ORF exon (Figure 2), for the same tissue RNA samples assayed with the intron-spanning set. The NAT2 ORF assay results showed a somewhat poorer signal-to-noise ratio, as indicated by a general tendency to higher Ct values, and the presence of genomic DNA contamination interfered with measurements of a few samples (Figure 4B). The detected hierarchy of tissue-specific NAT2 mRNA expression was very similar for the intron-spanning and the ORF TaqMan assays in the tissue RNA panels from Ambion (Figures 3A and 3B) and BD (Figures 4A and 4B). Thus, there was no evidence for any alternative NAT2 promoter(s) expressing NAT2 mRNA with a different tissue-specific profile nor was their evidence of sufficient quantity to substantially alter the relative level of NAT2 mRNA established by transcription at the -8.7 kb promoter locus.
Quantitative RT-PCR analysis of *NAT2* mRNA in human cell lines. Figure 5 shows *NAT2* TaqMan measurements for five tumor cell lines from breast (MCF-7), colon (Caco-2, SW480, and HT-29), liver (HepG2) and lung (A549) and a virally transformed embryonic kidney cell line (HEK293). None of the lines, including those derived from liver or colon, showed a level of *NAT2* expression that corresponded to that of the high-expressing normal tissues. The two cell lines with the highest *NAT2* expression, HepG2 and Caco-2, exhibited only about 1% and 5-10% respectively, of the *NAT2* expression in the corresponding tissue.
DISCUSSION

Accurate measurement of \textit{NAT2} expression in different human tissues is an important aspect of assessing its role in drug toxicity or carcinogenesis at various organ sites. In this study, a \textit{NAT2}-specific q-RT-PCR TaqMan assay with intron-spanning primers was designed based on the structure of \textit{NAT2} cDNAs in public databases and the design was validated by an extensive 5’-RLM-RACE analysis of \textit{NAT2} mRNA in liver. The accuracy of the assay for assessing relative mRNA levels in diverse tissues was verified by comparison to an alternative q-RT-PCR assay based on primers within the \textit{NAT2} open reading frame.

A recent study suggested that \textit{NAT2} transcripts also may be initiated immediately 5’ to the ORF-containing exon based on detection of RT-PCR products corresponding to such transcripts, most prominently in liver (Boukouvala and Sim, 2005). We have detected similar products by RT-PCR of liver and placental RNA, but RT-dependent PCR products of equal or greater prominence were also found using primer pairs from entirely within the 8.7 kb intron or from segments including the \textit{NAT2} promoter exon and immediately adjacent intronic sequences (data not shown). These results suggested that unspliced or incomplete primary \textit{NAT2} transcripts from the -8.7 kb promoter are present in RNA from liver and other tissues and that functional activity of an ORF-adjacent promoter cannot be directly inferred from demonstration of RT-PCR products derived from that region. The location of any \textit{NAT2} promoter immediately 5’ to the \textit{NAT2} ORF also is inconsistent with the 5’ structure of cDNAs in the public databases and our finding that 0/41 of the \textit{NAT2} cDNAs characterized by 5’-RLM-RACE of liver mRNAs had a 5’
terminus at that position. Together, the findings that NAT2 transcripts in liver originate predominantly at the -8.7 kb promoter and that both the intron-spanning (Figures 3A and 4A) and ORF-only (Figures 3B and 4B) TaqMan assays show similar relative levels of NAT2 mRNA in liver and other tissues suggest that any other possible alternative promoter in the non-hepatic tissues must be relatively very weak. These results also suggest that the pattern of NAT2 transcription start sites at the -8.7 kb promoter in tissues other than liver is not substantially shifted to a location 3’ to the forward primer used for the q-RT-PCR assay, since such a pattern of transcription initiation would also be detected as a discrepancy between the reported intron-spanning and ORF-only measurements. Thus, in the absence of evidence for any alternative NAT2 transcription start site or splicing pattern, the intron-spanning TaqMan assay design shown in Figure 2 appears to be superior for comparing NAT2 mRNA levels in different human tissue samples because of its high sensitivity and the elimination of measurement errors caused by contaminating genomic DNA.

The reliability of the NAT2 measurements is also dependent on specificity of the assay which is important to verify empirically because of the high homology of the NAT2 and NAT1 sequences in the regions that include the TaqMan assay primers. The MCF-7 cell line is known to express a high level of NAT1 mRNA (Husain et al., 2004), but only minimal levels of NAT2 mRNA were detected in MCF-7 with the intron-spanning assay (Figure 5) or the ORF assay (data not shown), demonstrating that each assay is highly specific for NAT2.
None of the cell lines studied here, including ones derived from liver and colon, was found to have the high level of \( \text{NAT2} \) mRNA expression characteristic of those tissues. This finding is consistent with previous reports of very low or undetectable \( \text{NAT2} \) enzyme activity in liver-derived cell lines (Coroneos and Sim, 1993). The observed differences between human tissues and cultured immortalized cell lines are not unexpected, since gene expression patterns in transformed and cancer cell lines do not always reflect the tissue from which they are derived and can vary with culture conditions.

For the 45 human tissue RNA samples representing 29 different organ types, \( \text{NAT2} \) mRNA levels (Figures 3A and 4A) were highest in liver, with slightly lower amounts in small intestine and colon. The expression of \( \text{NAT2} \) in liver has long been recognized as the primary determinant of the human acetylator phenotype (Grant et al., 1990; Bendriss et al., 1998) and \( \text{NAT2} \) enzyme is most readily detected in liver, colon and intestine (Turesky et al., 1991; Sim et al., 2000). \( \text{NAT2} \) mRNA was also readily detected in both tested samples of kidney, lung, spleen and testis and single samples from fetal liver, spinal cord, stomach and uterus at a level about 1% compared to liver. \( \text{NAT2} \) mRNA was also detected in heart, skeletal muscle, thymus and trachea, albeit at 10-fold lower levels. Wide-spread expression of \( \text{NAT2} \) in human tissues has been previously reported (Windmill et al., 2000). The low level of \( \text{NAT2} \) mRNA in fetal liver compared to adult liver suggests that \( \text{NAT2} \) expression in humans may follow a similar developmental pattern as in mice (McQueen and Chau, 2003).
NAT2 mRNA expression in bladder is of particular interest, because very strong epidemiological associations have been established between the NAT2 slow acetylation phenotype and increased risk of cancer of the urinary bladder caused by arylamines (Garcia-Closas et al., 2005; Carreon et al., 2006; Hein, 2006). The NAT2 rapid acetylation phenotype has also often been associated with increased risk of colorectal cancer (Lilla et al., 2006; Moslehi et al., 2006). Although NAT2 mRNA was relatively high in colon, the NAT2 mRNA level in bladder was very low compared to other tested tissues, about 0.1% of the liver level. This result suggests caution in assuming that a gene modifying cancer susceptibility is more likely to be highly expressed in the affected tissue. The association of urinary bladder cancer with NAT2 genetic polymorphism could be due to impaired detoxification of carcinogens in the liver, where NAT2 mRNA expression is high. Alternatively, factors such as the concentrations of pro-carcinogens, the presence of competing or cooperating enzyme activities or other cell or organ-specific features could magnify the effects of NAT2 enzyme activity in the bladder. A third possibility is that mRNA levels and enzyme activities measured with extracts from whole organs do not reflect the activity of the specific population of cells in which the carcinogenic mutations arise.

The identification of the precise location of NAT2 transcription start sites by 5'-RLM-RACE permits a detailed comparison of the corresponding NAT1 P1 (Husain et al., 2004) and NAT2 promoter sequences. In the immediate vicinity of the NAT2 TSSs, there is 85% nucleotide identity with the NAT1 P1 promoter (Figure 1), similar to the 87% identity found in the coding region of these paralogous genes. Within this region, the
spectrum of NAT2 TSSs in liver is relatively limited, with 34/40 of the 5’-termini located at one of two closely spaced adenine residues. In contrast, the spectrum of NAT1 TSSs, previously determined in RNA from breast and the MCF-7 cell line (Husain et al., 2004) is more diffuse, with only a partial overlap with the location of the NAT2 TSSs (Figure 1). It is notable that the NAT1 P1 promoter is actively transcribed in essentially all human tissues (Barker et al., 2006) whereas high level NAT2 expression is limited to very few tissues. The more limited spectrum of NAT2 transcription initiation sites may thus reflect a transcription initiation complex that has a greater degree of structural definition due to the required binding of tissue-specific transcription factors. Another distinction of the NAT2 promoter that likely contributes to its different expression profile is the absence of the consensus Sp1 site that occurs in the NAT1 P1 promoter immediately upstream of the sequence shown in Figure 1.

In conclusion, NAT2-specific transcription start sites were identified. A sensitive and specific TaqMan RT-PCR assay with intron-spanning primers was developed and used, together with a second TaqMan RT-PCR assay based on amplification of a NAT2 ORF exon segment, to measure NAT2 mRNA in 29 different human tissues. NAT2 mRNA was highest in liver, small intestine and colon and readily detected in most other tissues albeit at levels 1% or less than in liver. Expression of NAT2 in various human tissues may be relevant to associations between NAT2 genetic polymorphism and drug toxicity or arylamine-induced cancers at these sites.
REFERENCES


promoters that generate different 5'-UTR splice variants with altered translational activities. Biochem J 387:119-127.


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Footnotes

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FIGURE LEGENDS

FIG 1. Summary of 5’ termini of \textit{NAT2} cDNAs from 5’-RLM-RACE analysis of liver RNA samples and from public databases with comparison to \textit{NAT1}. A) 5’ termini determined by 5’-RLM-RACE of liver samples from Ambion and BD respectively indicated by “a” and “b”. The positions of the most frequent TSS locations at -59 and -64 are located 8711 and 8716 nucleotides upstream of the \textit{NAT2} ORF. B) The 5’ termini of NCBI database clones from liver (D90042, BG569293, AV658623, AV658656, BG533459, BG563731, BG568400, BG618195, BG569272, BC015878, CB161982, CB161860, D90040); colon adenocarcinoma (BX095770, AI792606) and hepatocellular carcinoma (AV684197) are shown as X and the 5’ termini of the small intestine oligo-capped clones HSI05750, HSI08034 and HSI15615 from DBTSS are shown as O. C) \textit{NAT1} mRNA termini previously determined by 5’-RLM-RACE with RNA samples from breast and the MCF-7 cell line (Husain et al., 2004) are indicated by B and M respectively. The sequence corresponding to the intron-spanning forward q-RT-PCR primer is shown in italics. In the \textit{NAT1} DNA sequence, nucleotides not identical to the \textit{NAT2} sequence are shown in bold.

FIG. 2. Structure of the typical \textit{NAT2} mRNA in relation to the \textit{NAT2} gene and the relative locations of primers and probes used for the intron-spanning and ORF TaqMan q-RT-PCRs. The sequence of the primer marked IS-for is shown in italics in the \textit{NAT2} sequence shown in Figure 1.
FIG. 3. TaqMan q-RT-PCR analyses of NAT2 in human tissue RNAs from Ambion. Duplicate measurements are indicated by light and dark gray bars. A) Relative NAT2 mRNA expression determined by TaqMan with the intron-spanning assay. B) Relative NAT2 mRNA expression determined by TaqMan with the NAT2 ORF assay. An * marks tissues present in both tested panels.

FIG. 4. TaqMan q-RT-PCR analyses of NAT2 in human tissue RNAs from BD, a breast RNA sample from Ambion and the liver sample from Ambion included in the 5′-RACE analysis. Duplicate measurements are indicated by light and dark gray bars. A) Relative NAT2 mRNA expression determined by TaqMan with the intron-spanning assay. B) Relative NAT2 mRNA expression determined by TaqMan with the ORF assay. In panel B, measurements of heart, kidney, salivary gland, testis and thymus are omitted because reactions without addition of reverse transcriptase indicated the presence of DNA contamination sufficient to interfere with the ORF assay. An * marks tissues present in both tested panels.

FIG. 5. TaqMan q-RT-PCR analysis of NAT2, with the intron-spanning assay, in RNA extracted from human cell lines derived from liver (HepG2), colon (HT-29, SW480, Caco-2), lung (A549), and breast (MCF-7) tumors and embryonic kidney (HEK293). Duplicate measurements are indicated by light and dark gray bars. Relative NAT2 mRNA expression is shown on the same scale as Figure 4A, with comparison to kidney, spleen and testis measurements also shown in Figure 4A.
Figure 1

A

B

C

TTCCACTGACATCACTTCCTGCAAGACCTTGGAGGGAGACGACTTATTTATAGACCTTTGGAAAGGCAAGGATCTGCATTCAGGCTAGTCCCTGGTTGCTGGCCAAAAGTAACT

TTCCACTGACATCACTTCCTGCAAGACCTTGGAGGGAGACGACTTATTTATAGACCTTTGGAAAGGCAAGGATCTGCATTCAGGCTAGTCCCTGGTTGCTGGCCAAAAGTAACT

TTCCARTGAGACACTTCCCTGCAAGACCTTGGAGGGAGACGACTTATTTATAGACCTTTGGAAAGGCAAGGATCTGCATTCAGGCTAGTCCCTGGTTGCTGGCCAAAAGTAACT

splice

splice

splice

15nt's CTAACT
Figure 2

Gene

mRNA

TSS region

ORF exon

<102

8,658

>870

IS-for IS-pr

IS-rev

ORF-for ORF-pr

ORF-rev
Figure 3A

Ambion series with NAT2 intron-spanning assay

Relative mRNA expression

Adipose
Bladder*
Brain*
Cervix
Colon*
Esophagus
Heart*
Kidney*
Liver*
Lung*
Ovary
Placenta*
Prostate*
Skel. muscle*
Spleen*
Testis*
Thymus*
Thyroid
Trachea*
Liver (BD)

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Figure 3B

Ambion series with NAT2 ORF assay

Relative mRNA expression

- Adipose
- Bladder
- Brain
- Cervix
- Colon
- Esophagus
- Heart
- Kidney
- Liver
- Lung
- Ovary
- Placenta
- Prostate
- Sm. intestine
- Spleen
- Testis
- Thymus
- Thyroid
- Trachea
- Liver (BD)

0.001
0.01
0.1
1
10
100

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Figure 4A

BD series with NAT2 intron-spanning assay

Relative mRNA expression

0.001 0.01 0.1 1 10 100

Adrenal Gland  | Bladder *  | Brain *  | Cerebellum  | Colon *  | Fetal Brain  | Fetal Liver | Kidney *  | Liver *  | Lung *  | Spleen *  | Stomach  | Spinal Cord  | Thymus *  | Trachea *  | Uterus  |

Breast  | Brain  | Cerebellum  | Fetal Brain  | Fetal Liver | Kidney *  | Liver *  | Liver (5'-RACE)  | Liver  | Sm. intestine *  | Sm. intestine *  | Stomach  | Stomach  | Testis *  | Testis  | Thymus *  | Thymus  |

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Figure 4B

BD series with NAT2 ORF assay

Relative mRNA expression

Adrenal Gland  Bladder *  Breast  Brain  *  Cerebellum  Colon  *  Fetal Brain  Fetal Liver  Lung  *  Placenta  *  Prostate  *  Spleen  *  Stomach  Sm. intestine  *  Spinal Cord  Trachea  *  Uterus  Liver (5'-RACE)  Liver (5'-RACE)
Figure 5

Cell lines with NAT2 intron-spanning assay

- Kidney
- Spleen
- Testis
- HepG2 #1
- HepG2 #2
- HT-29
- Caco-2
- A549
- HEK293

Relative mRNA expression