Human Arylamine N-Acetyltransferase 1 (NAT1) Knockout in MDA-MB-231 Breast Cancer Cell Lines Leads to Transcription of NAT2

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Many cancers, including breast cancer, have shown differential expression of human arylamine N-acetyltransferase 1 (NAT1). The exact effect this differential expression has on disease risk and progression remains unclear. While NAT1 is classically defined as a xenobiotic metabolizing enzyme, other functions and roles in endogenous metabolism have recently been described providing additional impetus for investigating the effects of varying levels of NAT1 on global gene expression. Our objective is to further evaluate the role of NAT1 in breast cancer by determining the effect of NAT1 overexpression, knockdown, and knockout on global gene expression in MDA-MB-231 cell lines. RNA-seq was utilized to interrogate differential gene expression (genes correlated with NAT1 activity) across three biological replicates of previously constructed and characterized MDA-MB-231 breast cancer cell lines expressing parental (Scrambled), increased (Up), decreased (Down, CRISPR 2–12), or knockout (CRISPR 2–19, CRISPR 5–50) levels of NAT1. 3,889 genes were significantly associated with the NAT1 N-acetylation activity of the cell lines (adjusted p ≤ 0.05); of those 3,889 genes, 1,756 were positively associated with NAT1 N-acetylation activity and 2,133 were negatively associated with NAT1 N-acetylation activity. An enrichment of genes involved in cell adhesion was observed. Additionally, human arylamine N-acetyltransferase 2 (NAT2) transcripts were observed in the complete NAT1 knockout cell lines (CRISPR 2–19 and CRISPR 5–50). This study provides further evidence that NAT1 functions as more than just a drug metabolizing enzyme given the observation that differences in NAT1 activity have significant impacts on global gene expression. Additionally, our data suggests the knockout of NAT1 results in transcription of its isozyme NAT2.

Keywords: NAT1, arylamine N-acetyltransferase 1, RNA-seq, NAT2, arylamine N-acetyltransferase 2, cell adhesion, breast cancer
1 INTRODUCTION

It is estimated by the American Cancer Society that breast cancer will account for 30% of new cancer cases and 15% of cancer-related deaths in women in 2021 (Siegel et al., 2021). When breast cancer is detected at the localized stage the 5-year survival rate is 99% however when detected at the distant stage the 5-year survival rate is a disappointing 28% (Siegel et al., 2018). These statistics highlight the need for a better understanding of breast cancer risk and progression so that interventions can be provided at the early stage when breast cancer is the most treatable and to identify novel molecular targets for treatment.

Human arylamine N-acetyltransferase 1 (NAT1) has been extensively studied in breast cancer. NAT1 is a classically described phase II xenobiotic metabolizing enzyme, but more recently has been shown to have additional, orthogonal, and diverse roles in metabolism including the hydrolysis of acetyl-Coenzyme A (acetyl-CoA) (Laurieri et al., 2014; Stepp et al., 2015; recently has been shown to have additional, orthogonal, and described phase II xenobiotic metabolizing enzyme, but more extensively studied in breast cancer. NAT1 is a classically described phase II xenobiotic metabolizing enzyme, but more

2 MATERIALS AND METHODS

2.1 Description of Cell Line Samples

Six cell lines constructed via both siRNA and CRISPR/Cas9 technologies from the MDA-MB-231 triple negative breast cancer cell line to have varying levels of human NAT1 N-acetylation activity were utilized in this study: Scrambled (transfection control with parental MDA-MB-231 activity; 9.41 ± 1.15 nmoles acetylated p-aminobenzoic acid/min/mg protein), Up (stable integration of plasmid overexpressing NAT1 with increased activity; 63.5 ± 10.3 nmoles acetylated p-aminobenzoic acid/min/mg protein), Down (stable transfection with siRNA specific to NAT1 with decreased activity; 6.55 ± 0.826 nmoles acetylated p-aminobenzoic acid/min/mg protein), CRISPR 2–12 (single allele deletion with CRISPR/Cas9 guide RNA 2 with decreased activity; 5.33 ± 0.572 nmoles acetylated p-aminobenzoic acid/min/mg protein), CRISPR 2–19 (double allele deletion with CRISPR/Cas9 guide RNA 2 with complete knockout of NAT1; no detectable activity), and CRISPR 5–50 (double allele deletion with CRISPR/Cas9 guide RNA 5 with complete knockout of NAT1; no detectable activity). Details on the construction and characterization of these cell lines has been described in detail elsewhere (Carlisle et al., 2018; Stepp et al., 2019). The Scrambled, Up, and Down cell lines are non-clonal cell lines while the CRISPR 2–12, CRISPR 2–19, and CRISPR 5–50 cell lines were established from a single isolated clone following transfection. Additionally, we acknowledge that the Scrambled cell line may have changes in gene expression not present in the other cell lines if the randomly generated scrambled shRNA utilized was complementary to anywhere in the genome. This risk was minimized by the chosen design/sequence of the shRNA.

2.2 Collection of Samples

Figure 1 illustrates the experimental approach and workflow of sample collection described below. Cells were plated in triplicate per biological replicate at a density of 500,000 cells per 150 mm × 25 mm cell plate for both transcriptomics and metabolomics analysis. All cell lines were cultured in high-glucose Dulbecco’s Modified Eagle Medium (DMEM), with 10% fetal bovine serum, 5% glutamine, and 5% penicillin/streptomycin added. Cells were allowed to grow for 3 days at 37°C and 5% CO₂ in an incubator.

Cells were then harvested on ice by adding 5 ml 0.25% trypsin and scraping the cells from the plate. Three plates were combined to form one sample (biological replicate). Three biological replicates for each cell line were collected for transcriptomics analysis. After harvesting the cells, cells were washed 3 times with ice-cold 1 x PBS. All supernatant was removed and 100 µl of cell pellet was reserved for transcriptomics analysis. The remaining cell pellet was reserved for metabolomics analysis. Samples were then flash frozen by placing in a pool of liquid nitrogen for 1 min and stored in an −80°C freezer until RNA isolation.

2.3 Transcriptomics

2.3.1 RNA Isolation

Total RNA was isolated from MDA-MB-231 breast cancer cells expressing parental (Scrambled), increased (Up), decreased (Down, CRISPR 2–12), and knockout (CRISPR 2–19, CRISPR 5–50) levels of NAT1 using the RNeasy® Mini Kit (Qiagen Sciences, Germantown, Maryland) according to manufacturer’s instructions. RNA quality was evaluated and concentrations were measured in each sample using a NanoDrop Bioanalyzer (Thermo Fischer Scientific).

2.3.2 Library Preparation

Libraries were prepared using the TruSeq Stranded mRNA LT Sample Prep Kit- Set A (Illumina, San Diego, California; Cat# RS-122-2101) with poly-A enrichment per manufacturer’s instructions. One µg of total RNA (in a volume of 50 µl) from
each sample was used in library preparation. Briefly, the total RNA was fragmented to improve sequence coverage over the transcriptome. Next, the first strand of cDNA was synthesized from the cleaved RNA fragments that were primed with random hexamers using reverse transcriptase and random primers. Then the second strand of cDNA was synthesized, thus giving us double stranded blunt end cDNA. Next, a single ‘A’ nucleotide was added to the 3’ ends of the blunt fragments to prevent them from ligating to one another during the adapter ligation reaction. A corresponding single ‘T’ nucleotide on the 3’ end of the adapter provides a complementary overhang for ligating the adapter to the fragment. Then, multiple indexing adapters were ligated to the ends of the double stranded cDNA, preparing them for hybridization onto a flow cell. Next the DNA fragments were enriched using PCR to selectively enrich those DNA fragments that have adapter molecules on both ends and to amplify the amount of DNA in the library. The PCR was performed with a PCR Primer Cocktail that anneals to the ends of the adapters. Finally, 30 µl of eluted library was collected and stored at −20°C. To avoid skewing the representation of the library, the number of PCR cycles are minimized. This kit includes steps to validate and normalize constructed libraries and methods to check quality control.

2.3.3 Library Validation

Quality: Size, purity and semi quantitation were performed on an Agilent Bioanalyzer using the Agilent DNA 1000 Kit. The final fragment size for all the samples was approximately 300bp which is expected according to the protocol.

Quantity: Sequencing library quantitation was performed by qPCR using the KAPA Library Quantitation Kit for Illumina Platforms. The standard curve method was used for quantitation using one to five DNA standards that came with the kit.

Normalize and Pool Libraries: Ten µl of sample was transferred from the wells to a new MIDI plate. We then normalized the concentration of the libraries to 10 nM using Tris-HCl 10 mM, pH 8.5 with 0.1% Tween 20. Five µl of each sample was then transferred to be pooled into a new LowBind 1.5 ml micro centrifuge tube for tube of 60 µl pooled 10 nM library. Then 4 nM dilution was made from the 10 nM pooled library by diluting with Tris-HCl 10 mM, pH 8.5 with 0.1% Tween 20.

Denaturing and diluting Libraries for the Nextseq 500: A total volume of 1.3 ml of 1.8 pM denatured library is needed for sequencing using the v2 kit. Poled 4 nM library was denatured by mixing with diluted NaOH and incubated at room temperature for 5 min 200 mM Tris HCl, pH 7.0 was then added. The reaction mixture was diluted to 20 pM using a pre-chilled Hybridization Buffer. 20 pM denatured library was further diluted to 1.8 pM using the same Hybridization Buffer. Before loading onto the reagent cartridge, 1.3 µl of denatured 20 pM Phix control was added to the 1,299 µl of denatured 1.8 pM library to a total volume of 1.3 ml for the first sequencing run (for the 2nd sequencing run 1.9 pM library was used).

2.3.4 Sequencing

Sequencing was performed on the University of Louisville Center for Genetics and Molecular Medicine’s (CGeMM) Illumina NextSeq 500 using the NextSeq 500/550 75 cycle High Output Kit v2 (FC-404-2005). A second run was performed to increase the number of reads. For each run, 72 single-end raw sequencing files (.fastq) (Cock et al., 2010) representing six conditions with three biological replicates and four lanes per replicate were generated. Reads were 75 base pair, single-end reads.

2.3.5 RNA-Seq Analysis

Adapter sequences were trimmed from resulting reads using Trim Galore (version 0.6.6). Quality of the sequencing reads (after trimming) were interrogated with FastQC (Andrews, 2014) (version 0.11.9) and found to be excellent. Reads were then aligned to the GENCODE release 31 GRCh38.p12 genome using the GENCODE v31 transcript annotation with the STAR (Dobin et al., 2013) aligner (version 2.7.6) using basic 2-pass mapping, with all 1st pass junctions inserted into the genome indices on the fly. Supplementary Table S1 indicates the number of reads successfully aligned for each of the samples.

Analysis and visualization of the resulting data were performed using R version 4.0.3 (R Foundation for Statistical Computing, Vienna, Austria). Aligned reads were quantified using the GenomicAlignments (Lawrence et al., 2013) package (Bioconductor) with the “IntersectionStrict” setting. Resulting read counts were log base two transformed and normalized using the rlog function of DESeq2 (Love et al., 2014). Additionally, DESeq2 was utilized to estimate linear regression models relating gene expression and NAT1 N-acetylation activity. This method allows us to identify genes whose expression is significantly associated with NAT1 N-acetylation activity. Significance of linear regression analysis is reported as adjusted (for multiple comparisons) Wald test p-values. Principal component analysis (PCA) was conducted by singular value decomposition of the centered data matrix on the most 1,000 variable genes in the entire dataset. The scores of the first (x-axis) and second (y-axis) principal component were plotted.

Log2 change in gene expression per unit change in NAT1 N-acetylation activity and significance (adjusted p-values) from regression analysis were visualized using volcano plots. Unbiased clustering of samples by hierarchical clustering was conducted using the WPGMA method using the most significant 100 differentially expressed genes; resulting data were plotted as heatmaps for visualization. Pathway enrichment analysis was conducted utilizing the linear regression results as input with the absolute value of the Wald test statistic as the variable for ordering. We utilized the normalized enrichment score to determine the relative degree of enrichment. Figure 1 illustrates the overall experimental approach and data analyses methods.

2.4 Quantitative Measurement of NAT1 and NAT2 mRNA

Quantitative Real-Time Reverse Transcription Polymerase Chain Reaction (qRT-PCR) was conducted for NAT1 and arylamine N-acetyltransferase 2 (NAT2) in each of the six constructed cell lines as previously described (Husain et al., 2007a; Husain et al., 2007b; Millner et al., 2012). Briefly, total RNA was isolated from
each cell line using the RNAeasy Mini Kit (Qiagen, Germantown, MD). Isolated RNA was used to transcribe cDNA using the High Capacity Reverse Transcriptase kit (Life Technologies, Carlsbad, CA). Resulting cDNA was utilized for quantitative measurement of NAT1 and NAT2 via qRT-PCR. TaqMan analysis was performed using the ABI 7700 sequence detection system (Applied Biosystems, Foster City, CA). The gene probes utilized were designed previously to discriminate between NAT1 and NAT2. Student’s t-tests were utilized for testing significance.

### 2.5 NAT2 N-Acetylation Activity Assays

*In vitro* NAT2 N-acetylation activity was determined in each constructed cell line as previously described (Grant et al., 1991; Hein et al., 1993). Briefly, cell lysate from each cell line was incubated with 1 mM acetyl-coenzyme A and 300 μM sulfamethazine (SMZ) at 37°C for 10 min. Reactions were terminated with the addition of 1/10 reaction volume 1 M acetic acid. Reaction products were collected and analyzed using an Agilent Technologies 1,260 Infinity high performance liquid chromatography (HPLC) using a LiChrospher 100 RP-18 (5 μm) column to determine the amount of acetylated product. Our reported limit of detection for N-acetylated SMZ is 0.15 nmol/min/mg.

### 3 RESULTS

PABA N-acetylation activity was measured in constructed MDA-MB-231 cell lines and reported previously (Carlisle et al., 2020). Briefly, the Scrambled cell line had approximately the same activity as the Parent MDA-MB-231 cell line with no manipulation while the Up cell line had an approximate 700% increase in activity. Additionally, the Down and CRISPR 2–12 cell lines had approximately 65 and 50% of the activity of the Parent and Scrambled cell lines, respectively, while the CRISPR 2–19 and CRISPR 5–50 cell lines had no detectable activity.

On a per-sample average, 91% of reads aligned to the human genome resulting in approximately 34.6 M uniquely aligned reads. Regression analysis revealed that the expression of 3,889 genes were significantly associated with the NAT1 N-acetylation activity of the cell lines (*adjusted p* ≤ 0.05); of those 3,889 genes, 1,756 were positively associated with NAT1 N-acetylation activity and 2,133 were negatively associated with NAT1 N-acetylation activity (Figure 2).

Principal component analysis of the 1,000 most variable genes in the dataset revealed that all cell lines in this study had distinct transcriptomic profiles as each clustered independently by group membership (Figure 3). In our dataset, principal component 1 explains 52.2% of the variance in the data while principal component 2 explains 20.8% of the variance. The CRISPR
2–12, CRISPR 2–19, and CRISPR 5–50 groups are separated from the other three groups along principal component 1. The CRISPR 2–12 and CRISPR 5–50 cell lines were additionally separated from the remaining 4 cell lines along principal component 2. This revealed that the Scrambled, Up, and Down groups had transcriptomic profiles similar to each other but different from the other three groups suggesting the construction method utilized may have led to unique differences.

Unsupervised hierarchical clustering of the 100 most significant genes associated with NAT1 N-acetylation activity was conducted to visualize gene expression patterns within all samples and to identify groups of genes with similar expression patterns (Figure 4). We observed one cluster of 24 genes, that included NAT1, to be positively associated with NAT1 N-acetylation activity. Gene expression in this cluster was generally low in the CRISPR 2–19 and CRISPR 5–50 cell lines, higher in the Down, CRISPR 2–12, and Scrambled cell line, and highest in the Up cell line. The remaining 76 genes which grouped as a separate cluster, exhibited the opposite pattern; gene expression in this cluster was generally highest in the Up cell line. The remaining 76 genes which grouped as a separate cluster, exhibited the opposite pattern; gene expression in this cluster was generally highest in the Up cell line. The remaining 76 genes which grouped as a separate cluster, exhibited the opposite pattern; gene expression in this cluster was generally highest in the Up cell line.

Gene encoding proteins with functions related to acyl/acyetyl transfer and binding were significantly associated with NAT1 N-acetylation activity. Vesicle amine transport 1 like (VAT1L) which is involved in acyl group transfer, was inversely related to NAT1 activity. Acyl-CoA binding domain containing 7 (ACBD7), was positively associated with NAT1 N-acetylation activity. Glucosaminyl (N-acetyl) transferase 4 (GCNT4) was negatively associated with NAT1 N-acetylation activity. Additionally, a negative association between 5-hydroxytryptamine receptor 1F (HTR1F) and NAT1 N-acetylation activity was observed.

Many protocadherin and cadherin genes were found to be significantly associated with NAT1 activity in the cell lines (Supplemental Table S2). Twenty-nine protocadherin genes were significantly associated with NAT1 activity (with no significant association observed in 33) and all but five of those genes were associated with an increase in expression per unit NAT1 activity. These included alpha, beta, and gamma subfamilies. Five cadherin genes (CDH2, CDH3, CDH11, CDH13, CDH18) were significantly associated with NAT1 activity and expression of all five were decreased per unit of NAT1 activity. Additionally, three of the four FAT atypical cadherin genes (FAT1, FAT3, FAT4) were significantly associated with NAT1 activity however the fold-changes were very small per NAT1 activity unit and in different directions; FAT1 and FAT4 were increased with NAT1 activity while FAT3 was decreased with NAT1 activity. Transmembrane O-mannosyltransferase targeting cadherins 1 (TMTC1) was significantly associated with NAT1 activity and expression was decreased per unit NAT1 activity.

3.1 NAT Isozyme Expression in Transcriptomics Dataset

The expression of NAT1 and its isozyme, NAT2, were compared between all cell lines. As expected, given the N-acetylation activities
The 100 most differentially expressed genes cluster into two overall groups with distinguishable expression profiles. This analysis shows each cell line has a distinct expression profile for the 100 most significant genes associated with NAT1 activity and that there are many genes with similar expression profiles across the samples. There are two overall groups that form in terms of similar patterns of expression, those that are positively associated with NAT1 activity (24—top cluster) and those that are negatively associated with NAT1 activity (76—bottom cluster). Each row of the heatmap shows the expression for a single gene while each column shows gene expression for an individual sample. Expression values have been transformed to Z-scores with red colors representing high expression and blue colors representing low expression.
measured, NAT1 expression was decreased in the Down, CRISPR 2–12, CRISPR 2–19, and CRISPR 5–50 cell lines and increased in the Up cell line compared to the Scrambled cell line. Notably, in the two complete NAT1 knockout cell lines (CRISPR 2–19 and CRISPR 5–50), NAT2 expression was observed while the expression was much lower in the other cell lines (Figure 5). We visualized the RNA-seq reads that were mapping to NAT2 using the Integrated Genomics Viewer (IGV) to double check that the reads were not NAT1 reads mis-mapping to NAT2 given the isozymes high degree of sequence homology (Figure 6); we confirmed these were not NAT1 misreads as we observed known NAT2 SNPs in the mapped sequence reads. The reads mapping to NAT2 had the following SNPs within them: rs1801280 T > C and rs1799929 C > T which corresponds with the NAT2*5A haplotype. We also verified the observation of NAT2 reads with qRT-PCR analysis for NAT1 and NAT2 in all 6 cell lines (Figure 7). Although not significant for all cell lines, we observed a pattern of decreased NAT1 transcripts and increased NAT2 transcripts in samples. We conducted NAT2 activity assays with cell lysate from all six constructed cell lines using the prototypic NAT2 substrate sulfamethazine (SMZ) however no NAT2 activity was detected in any cell line (data not shown). Currently, the functionality of the NAT2 transcripts produced in the NAT1 knockout cell lines is unknown.

3.2 Pathway Analysis

Pathway enrichment analysis was conducted using Gene Set Enrichment and the KEGG pathways to determine what metabolic pathways were enriched for genes associated with NAT1 activity (Figure 8). We focused on pathways that had a normalized enrichment score of >1.40. The Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways were used (Ogata et al., 1999; Kanchisa and Goto, 2000). Disease associated pathways were removed from the analysis results. Many pathways were significantly enriched for differences including hematopoietic cell lineage, cell adhesion molecules, antigen processing and presentation, IL-17 signaling, intestinal immune network for IGA production, caffeine metabolism, and antifolate resistance pathways were found to be significantly enriched. This further supports our observation that NAT1 activity is strongly associated with the expression of cell adhesion genes. Additionally, pathways that we would expect to be affected by NAT1 given previous literature and knowledge, such as caffeine metabolism and antifolate resistance, were found to be significantly enriched for differences. Some of the enriched pathways additionally suggest a relationship between the immune system and NAT1.

4 DISCUSSION

As our samples for this study were genetically modified breast cancer cell lines that had been passaged for a few growth cycles, we decided to utilize a regression approach for data analysis rather than pair-wise comparisons to our control cell line. This approach decreases the likelihood that genes identified to be differentially expressed between groups are due to off-target effects of CRISPR/Cas9 editing and/or additional mutations incurred during passaging of cell lines or during clonal selection after CRISPR/Cas9. One limitation of this approach, however, is that it relies on NAT1 N-acetyltransferase activity being the functional measure for this enzyme’s activity and may limit us identifying genes that are affected if NAT1 has any enzyme functions independent of this activity. Nevertheless, we should still be able to detect some of those possible independent effects because we know our complete NAT1 knockout cell lines have mutations that lead to very truncated NAT1 protein that have lost nearly all protein domains. Additionally, it has been noted by others that CRISPR/Cas9 genetic editing frequently leads to on-target mRNA misregulation which leads to degradation of the mutant mRNA produced (Tuladhar et al., 2019).

In recent years many roles for NAT1 independent of N-acetylation of exogenous xenobiotics have been shown. One challenge of identifying all of the potential roles of NAT1 in
**FIGURE 6** | Sequencing Reads Map to the NAT2 Gene But Only in the Two Knockout Cell Lines, CRISPR 2–19 and CRISPR 5–50. The mapping of reads to the NAT2 gene were visualized using the Integrated Genomic Viewer in the Scrambled, CRISPR 2–19, and CRISPR 5–50 cell lines. Each linear gray bar represents an individual read. The red and blue lines represent positions in the read where a single nucleotide polymorphism compared to the reference genome is present. Zero to very few reads mapped to NAT2 in the Scrambled cell line samples, many reads mapped to NAT2 in the CRISPR 2–19 cell line samples, and a medium amount of reads mapped to NAT2 in the CRISPR 5–50 cell line samples.

**FIGURE 7** | RT-qPCR confirms inverse relationship between NAT1 and NAT2 expression in our cell lines. PCR probes designed to discriminate between NAT1 and NAT2 were utilized for real-time quantitative polymerase chain reactions. mRNA expression was normalized to the expression of the parental MDA-MB-231 cell line. A trend of decreased NAT1 expression and increased NAT2 expression was observed. *p ≤ 0.05, ***p ≤ 0.001.
metabolism has been the approaches taken thus far. When performing targeted assays probing NAT1’s effect on specific pathways/genes we are limited by what is known when deciding where to look. In this study we have taken a global systems biology approach which removes that previous limitation.

In this study we identified additional supportive evidence of an association between NAT1 and cell adhesion molecules. Many have noted that overexpression NAT1, specifically in breast cancer, is associated with increased metastasis (Smid et al., 2006; Savci-Heijnink et al., 2016; Zhao et al., 2020) and growth (Adam et al., 2003) and conversely knockdown or deletion of NAT1 leads to increased adherence to collagen (Li et al., 2020) and decreased invasion (Tiang et al., 2010, 2015; Li et al., 2020), E-cadherin up-regulation and cell-cell contact growth inhibition (Tiang et al., 2011), and reduced anchorage independent growth (Stepp et al., 2018; Stepp et al., 2019). Here we have shown the expression of many members of the cadherin family of proteins are associated with NAT1 N-acetylation activity. Particularly, the negative association between the genes CDH2, CDH3, CDH11, CDH13, CDH18 and NAT1 N-acetylation activity supports observations of high NAT1 expression in breast cancer and increased metastasis.

The cadherin family of genes are subject to numerous regulatory mechanisms (Gumbiner, 2000) complicating the interpretation of the relationship between cadherins and NAT1 activity directly. However, we additionally observed that expression of TMTC1, a gene that encodes a protein important for adding O-mannosyl post-translational modifications to members of the cadherin family, was inversely related to NAT1 activity; it has been shown by others that TMTC3, a closely related family member to TMTC1, is important for ensuring E-cadherin operates properly in cell adhesion (Graham et al., 2020); when TMTC1 was knocked out cells showed increased migration ability (Graham et al., 2020). Our pathway analysis indicated the KEGG pathway: cell adhesion molecules, was one of the most significantly enriched pathways for genes associated with NAT1 activity. Although the mechanism behind this relationship remains unclear, this adds additional evidence of a relationship and further impetus to study this phenomenon more closely in order to potentially elucidate NAT1 as an anti-metastasis drug target.

The observation that the complete knockout, but not knockdown, of NAT1 lead to NAT2 transcript production leads us to hypothesize that there may be a compensation mechanism that occurs when NAT1 N-acetylation is lost, suggesting NAT1 has an essential role. There have been examples of humans that have no detectable NAT1 N-acetylation activity (Bruhn et al., 1999) however NAT2 expression and/or activity has never been investigated specifically in this population. Notably, a retrospective analysis of publicly available NAT1 and NAT2 gene expression data in established breast cancer cell lines, primary breast tumor tissues, and normal breast tissues showed a small positive correlation between the two genes (Carlisle and Hein, 2018). One reason that study did not observe an inverse relationship between NAT1 and NAT2 gene expression may be because complete knockout of NAT1 may be necessary before the compensation mechanism occurs given we only observed NAT2 transcripts in the complete NAT1 KO cell lines but not the cell lines with decreased NAT1. Data reported by the Cancer Cell Line Encyclopedia (CCLE) indicates the MDA-MB-231 cell line is diploid for NAT2 and does not show copy number variation (Ghandi et al., 2019).

The relationship between NAT1 and NAT2 expression has not been studied in-depth although it is known each has tissue specific expression. NAT1 is expressed throughout most tissues while NAT2 is most highly expressed in the liver and digestive tract. Since NAT1 and NAT2 are isozymes with overlapping substrate specificities (although distinct) it has been thought that the isozyme with the highest expression would be most important to understand in a given tissue. While our RNA-seq data showed the highest levels of NAT2 transcripts in the CRISPR 2–19 cell line, our qRT-PCR showed the highest levels of NAT2 transcripts in the CRISPR 5–50 cell line. We hypothesize that this is because the samples for each set of experiments were collected at different times; the phase of cell growth as well as growth conditions may affect the mechanism responsible for transcription of NAT2. Additionally, others have noted that CRISPR/Cas9 genetic editing can lead to degradation of the mutant mRNAs produced (NAT1 in our experiments); we hypothesize that this process may also be affected by the growth conditions or growth phase during sample collection.

Additionally, the associations observed between NAT1 N-acetylation activity and other genes closely related to transfer of acyl groups such as, VAT1L (inversely related to NAT1 activity), GCNT4 (negatively associated with NAT1 N-acetylation activity), and ACBD7 (positively associated with NAT1 N-acetylation activity) suggest there are mechanisms occurring to maintain homeostasis as it relates to acetyl group transfers. ACBD7 is a member of a family of proteins involved in
numerous intracellular processes including fatty acid-, glycerolipid- and glycerophospholipid biosynthesis, β-oxidation, cellular differentiation and proliferation as well as in the regulation of numerous enzyme activities (Nees et al., 2015). This association may help us better understand the finding in our previous metabolomics study that the β-oxidation of fatty acids seemed to be impaired in NAT1 deficient cell lines (Carlisle et al., 2020).

As NAT1 is known to N-acetylate serotonin (Steinberg et al., 1969; Backlund et al., 2017), albeit at very low levels in comparison to aralkylamine N-acetyltransferase (AANAT), we find the negative association between 5-hydroxytryptamine receptor 1F (HTRIF) and NAT1 N-acetylation activity interesting. The negative association between HTRIF and NAT1 N-acetylation activity indicates an upregulation of the serotonin receptor when NAT activity is low and presumably less serotonin is N-acetylated and a downregulation of the serotonin receptor when NAT activity is high and presumably more serotonin is N-acetylated. In our previous metabolomics study of the same samples, a positive association was observed between NAT1 N-acetylation activity and serotonin metabolite abundance (Carlisle et al., 2020). Taken together, this study suggests there are complex homeostatic/compensatory mechanisms occurring based on NAT1 levels as well as other factors such as AANAT levels.

5 CONCLUSION

This is the first study to measure the global transcriptome profile of breast cancer cells with varying levels of NAT1 N-acetylation activity. The expression of many genes, especially genes in the cadherin family, were significantly associated with NAT1 N-acetylation activity suggesting either regulation by NAT1, regulation by acetyl-CoA (or products of its hydrolysis), co-regulation mechanisms, or mechanisms compensating for the loss of NAT1. The most notable finding of this study was the observation that NAT2 gene expression was significantly inversely associated with NAT1 N-acetylation activity and this relationship was also observed via RT-PCR methods. It appears only the complete knockout of NAT1 stimulates transcription of NAT2 suggesting complete loss of NAT1 N-acetylation activity is necessary. This observation suggests NAT2 transcripts are produced as a compensation mechanism for the complete loss of NAT1 however the mechanism by which the cell senses the loss of NAT1 and transcribes NAT2 has not been identified and requires further investigation. As the two NAT1 knockout cell lines (nor any of the other cell lines) did not have any detectable N-acetylation activity toward the prototypic NAT2 substrate SMZ (limit of detection = 0.15 nmol/min/mg), the functionality of the NAT2 transcripts remains unknown and warrants further investigation.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are publicly available. This data can be found here: https://www.ncbi.nlm.nih.gov/Traces/study/?acc=PRJNA774803.

AUTHOR CONTRIBUTIONS

SC and DH contributed to conception and design of the study. SC and MD prepared samples. SC analyzed the resulting data and prepared figures. PT assisted with statistical analysis of the data. SC wrote the first draft of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphar.2021.803254/full#supplementary-material

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