

ORIGINAL ARTICLE

Pharmacometabolomic Assessments of Atenolol and Hydrochlorothiazide Treatment Reveal Novel Drug Response Phenotypes

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Achieving hypertension (HTN) control and mitigating the adverse health effects associated with HTN continues to be a global challenge. Some individuals respond poorly to current HTN therapies, and mechanisms for response variation remain poorly understood. We used a nontargeted metabolomics approach (gas chromatography time-of-flight/mass spectrometry gas chromatography time-of-flight/mass spectrometry) measuring 489 metabolites to characterize metabolite signatures associated with treatment response to anti-HTN drugs, atenolol (ATEN), and hydrochlorothiazide (HCTZ), in white and black participants with uncomplicated HTN enrolled in the Pharmacogenomic Evaluation of Antihypertensive Responses study. Metabolite profiles were significantly different between races, and metabolite responses associated with home diastolic blood pressure (HDBP) response were identified. Metabolite pathway analyses identified gluconeogenesis, plasmalogen synthesis, and tryptophan metabolism increases in white participants treated with HCTZ ($P < 0.05$). Furthermore, we developed predictive models from metabolite signatures of HDBP treatment response ($P < 1 \times 10^{-5}$). As part of a quantitative systems pharmacology approach, the metabolites identified herein may serve as biomarkers for improving treatment decisions and elucidating mechanisms driving HTN treatment responses.

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Study Highlights

WHAT IS THE CURRENT KNOWLEDGE ON THE TOPIC? The adverse health effects of HTN have been well characterized, and the impact of ATEN on metabolite signatures has been previously investigated. • WHAT QUESTIONS DID THIS STUDY ADDRESS? How endogenous metabolites and biological pathways are impacted in subjects treated with either ATEN or HCTZ. • WHAT THIS STUDY ADDS TO OUR KNOWLEDGE Biological pathways indicating changes in gluconeogenesis and tryptophan metabolism were impacted upon treatment with HCTZ. Furthermore, multivariable models of metabolites are capable of predicting treatment response to patients administered ATEN or HCTZ. • HOW THIS MIGHT CHANGE CLINICAL PHARMACOLOGY AND THERAPEUTICS The work presented here is an early step toward identifying metabolite biomarkers for treatment response to hypertensive therapies.

Hypertension (HTN) remains a public health burden affecting about one-third of US adults and more than one billion individuals worldwide.^{1,2} Response to antihypertensive treatment also varies among individuals with some individuals remaining resistant to certain therapies, and only about 50% of patients treated with antihypertensive therapies achieve blood pressure (BP) control.^{1,3–5} Ethnic differences regarding both side effects and drug efficacy have been noted between white and black populations prescribed antihypertensives.^{3,5} For these reasons, the current approach for treatment of HTN is suboptimal. Thus, identifying predictors associated with BP response of antihypertensive therapies would be of value in optimizing treatment selection, and ultimately reduce morbidity and mortality by improving BP control. Significant contributions to the development of

more effective therapies can be made by developing a deeper understanding of the mechanism of action of current therapies, and mapping of pathways implicated in both treatment response and side effects.

Atenolol (ATEN) is a cardioselective beta-adrenergic receptor blocker that was previously a first-line treatment for HTN, but has more recently been downgraded because of its reduced efficacy at preventing adverse cardiovascular events compared with some other antihypertensive agents.^{3,6} Whether this reduced efficacy is due to ATEN's short duration of action and need for twice daily dosing, despite it typically being prescribed only once a day, is unclear. Clarifying the metabolic pathways impacted by ATEN treatment may help to identify which patients will respond favorably to ATEN treatment and can provide new

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insights for improved drug development. In a previous publication, we identified an initial metabolic signature of ATEN exposure.⁷ In this study, we expand upon these findings and construct a multivariable predictive model of home diastolic blood pressure (HDBP) response after ATEN treatment and interrogate pathways impacted upon ATEN treatment.

Hydrochlorothiazide (HCTZ) is a thiazide diuretic and is considered a first-line therapy for all populations.³ Additionally, HCTZ is largely well tolerated and considered comparably effective to many other treatment options, including angiotensin converting enzyme inhibitors, calcium channel blockers, and angiotensin receptor blockers.³ For black individuals, thiazide-based treatments are considered preferable, based on evidence that they offer superior prevention of cerebrovascular effects, heart failure, and other outcomes when compared to angiotensin converting enzyme inhibitor treatments.^{3,4} HCTZ is also widely available and inexpensive. However, there remains a significant portion of both black and white individuals that are resistant to HCTZ therapy, and, presently, no biomarkers exist that are capable of identifying which of these patients will experience adverse outcomes or which patients will be resistant to treatment. Identifying putative biomarkers and metabolic pathways implicated in this disparity of response is a goal of the present study.

Metabolomics, the study of metabolism at an “omic” level, integrates naturally with quantitative systems pharmacology approaches, and aims to improve our understanding of drug mechanisms of action and the molecular basis for drug response variation.⁸ Recently, several studies successfully used pharmacometabolomics to identify novel biomarkers and explain variation in drug response (e.g., sertraline, escitalopram, aspirin).^{8–14} Furthermore, metabolic phenotypes have been used to explain variation in BP across ethnic groups.¹³ In the present study, we used a nontargeted metabolomics approach in which we measured 489 metabolites (of both known and unknown annotation) in 206 and 227 participants treated with HCTZ and ATEN, respectively, to define metabolites significantly altered after exposure to either drug and significantly associated with change in HDBP. Furthermore, we performed exploratory analysis and constructed multivariable models predictive of changes in HDBP. Overall, through the systems-level metabolomics approach, our study provides new insights into the potential biological mechanisms of HTN and BP response in white and black individuals, and provides candidate metabolites as potential new biomarkers.

METHODS

Study population

Subjects were enrolled in the Pharmacogenomic Evaluation of Antihypertensive Responses study at the University of Florida (Gainesville, FL), Emory University (Atlanta, GA), and the Mayo Clinic (Rochester, MN). Pharmacogenomic Evaluation of Antihypertensive Responses is registered at clinicaltrials.gov, <http://clinicaltrials.gov/ct2/show/NCT00246519>. Enrolled participants were primary care patients of any self-identified race or ethnicity, aged 17–65 years, with newly diagnosed,

untreated, or treated, mild-to-moderate essential HTN. Plasma samples and associated clinical data were obtained from 128 white participants and 109 black participants randomly assigned to receive ATEN monotherapy, and 123 white participants and 83 black participants randomly assigned to receive HCTZ monotherapy. Individuals with cardiovascular disease, primary renal or hepatic disease, or diabetes mellitus (type I or II) were excluded from participating. Nineteen participants were taking statins before starting the trial or initiated statin treatment during the trial. However, statin was not incorporated into subsequent analyses because stratification of metabolic profiles by statin status was not statistically significantly different ($P > 0.1$). Additional details regarding the study population have been previously published in Johnson *et al.*¹⁵

Study protocol

The Pharmacogenomic Evaluation of Antihypertensive Responses protocol was approved by the institutional review boards at all study sites (University of Florida, Mayo Clinic, and Emory University) and after providing voluntary informed consent, all study participants were randomly assigned to either ATEN or HCTZ therapies. A minimum washout period of 18 days was initiated for any participant already receiving treatment for HTN. Upon completion of baseline measurements, 50 mg daily or 12.5 mg daily was initiated for ATEN or HCTZ groups, respectively. After three weeks of treatment, and based on BP $> 120/70$ mmHg and tolerability, subjects were titrated to 100 mg daily or 25 mg daily for ATEN and HCTZ groups, respectively. BP was recorded in triplicate upon rising from bed in the morning and before retiring in the evening. The morning and evening recorded triplicate BP measurements were subsequently averaged. BP measurements were obtained using a Microlife model 3AC1-PC home BP monitor (BP Microlife, Minneapolis, MN).¹⁵ In addition to baseline BP measurements, post-treatment BP was assessed after nine weeks of treatment. Baseline (before drug administration) and posttreatment plasma samples were collected during study visits under fasting conditions. Dietary information (e.g., food intake, food preference, sodium intake) were not recorded, although subjects were asked to maintain a steady diet during participation in the study, which minimizes the impact of dietary intake on the metabolite measurements. Additional information regarding the study design details have been previously published in Johnson *et al.*¹⁵

Metabolomic profiling

Study design information was entered into the miniX database (a simplified version of the SetupX database).¹⁶ All plasma samples were aliquoted and stored at -80°C until use, at which point 30 μL of each sample was thawed, extracted, and derivatized.¹⁷ Briefly, 30 μL aliquots were extracted with 1 mL of degassed acetonitrile:isopropanol:water (3:3:2) at -20°C , centrifuged, aliquoted into two portions, and evaporated to complete dryness. Acetonitrile/water (1:1) was used to remove membrane lipids and triglycerides and the supernatant was again dried down. Internal standards C8–C30 FAMES were added and the sample was derivatized using methoxyamine hydrochloride in pyridine and subsequently by MSTFA (Sigma-Aldrich) for trimethylsilylation of acidic

protons. All metabolites were measured as peak height. A total of 489 metabolites were measured (224 known and 265 unknown metabolites). Gas chromatography time-of-flight/mass spectrometry data acquisition and processing were conducted, as previously described.⁷

Data processing and statistical analysis

All data processing and analysis was performed in the open-source statistical software, R.¹⁸ Initial processing of the data included testing to determine if the peak height of each metabolite was normally distributed based on whether the data were skewed more or less upon natural log transformation. If data for a metabolite were skewed less upon natural log transformation, then the metabolite data was transformed and used for all subsequent analysis. Skewness was determined by the ratio of the number of metabolites that were $<1\sigma$ to the number of metabolites that were $>1\sigma$ from the mean (**Supplemental Figures S1 and S2**). Samples were considered outliers if ± 5 median absolute deviations from the median, and were subsequently removed from further analysis.

Signature of exposure

Metabolites significantly altered by treatment with either ATEN or HCTZ were determined using a pairwise Wilcoxon signed-rank test and testing both with and without stratifying by race. False discovery rate was used to correct for multiple comparisons.¹⁹ Metabolites with a $q < 0.2$ were considered to be statistically significant.

Signature of response

Comparisons of the number of participants of each race that did not respond to treatment with a decrease in HDBP were tested using a Fisher's exact test. Univariate linear regression analysis was used to test associations between HDBP and each covariate (baseline HDBP, baseline levels of renin, insulin, glucose, homeostasis model of assessment, triglyceride, uric acid, high-density lipoprotein, low-density lipoprotein, and race, gender, waist size (cm), age, and body mass index). Covariates associated with HDBP with $P < 0.05$ were considered to be statistically significant and were included in the linear metabolomics model to control for confounding. Subsequently, associations were performed using the initial/baseline metabolite level, post-treatment level, and the change observed between the initial and posttreatment level. If two covariates were highly correlated ($R^2 > 0.15$), then only the most significantly associated covariate was retained in the model. Associations were tested both with and without treatment stratification, and with and without race stratification.

Multivariable model

Before performing the multivariable associations, the dataset was partitioned into a discovery (50%) and validation set (50%) to internally assess potential predictive performance. Multivariable associations were determined by generating multiple regression models. Variable selection was performed by models with all combinations of the 15 most significant baseline metabolites and all significant covariates from the univariate analysis using the discovery data. Although significant metabolites were allowed to drop from

the individual models during variable selection, all significant covariates were maintained in each model iteration. The model with the lowest BIC was carried forward and tested in the validation dataset. Overall model performance was evaluated by a Bonferroni corrected P value and R^2 (with bootstrapped 95% confidence interval) using the validation data.

Hierarchical clustering

Significant metabolites for either white, black, or both groups treated with ATEN or HCTZ with $q < 0.2$ were used for clustering, respectively. Missing values were imputed with the median across metabolites for all other participants. Hierarchical clustering was performed using Modularized Modularity Clustering and Pearson correlation.²⁰

Pathway analysis

Metabolite lists were created for each treatment/race subgroup from the signature of exposure results. Groups were defined by drug (ATEN or HCTZ), direction of metabolite association (increased or decreased), and race. In order to combine metabolite level to a pathway level score, a correlated Lancaster approach²¹ was performed using data from the Human Metabolome Database version 3.5,²² and International Union of Pure and Applied Chemistry International Chemical Identifier codes for annotation. This approach uses correlation information from permutation testing of all known metabolites in the study rather than using a list based on a metabolite significance threshold. Pathway significance values were adjusted for multiple comparisons using a Bonferroni correction with a significance threshold of $P < 0.05$.

RESULTS

The vast majority of participants (83.1%) treated with either ATEN or HCTZ experienced a decrease in HDBP over the course of the nine-week treatment period, with an overall mean reduction in HDBP of -6.45 ± 0.16 mmHg (**Figure 1**). The white cohort treated with ATEN had the largest decrease in HDBP with a mean of -10.13 ± 0.6 mmHg. The black cohort treated with ATEN displayed the poorest response with a mean change in HDBP of -4.0 ± 0.64 mmHg and 77 (70.6%) participants recorded an overall decrease in HDBP. The white participants treated with HCTZ displayed a mean decrease in HDBP of -4.5 ± 0.51 mmHg with 102 of participants (83%) with an overall decrease in HDBP. Last, the black participants treated with HCTZ had a mean change in HDBP of -6.8 ± 0.73 mmHg, and 66 participants (79.5%) displayed an overall decrease in HDBP. Distributions of responses were statistically significantly different ($q < 0.05$, with matching letters) between ATEN and HCTZ across all groups when both combined and separated by race (**Figure 1**).

Signature of exposure to ATEN and HCTZ

The baseline metabolic profiles were significantly different between the black and white cohorts, based on a paired Wilcoxon rank test ($P < .001$), confirming the need to stratify the analyses by race. Additional information detailing our results from this analysis can be found in the supplementary material (**Supplementary Table S1**).

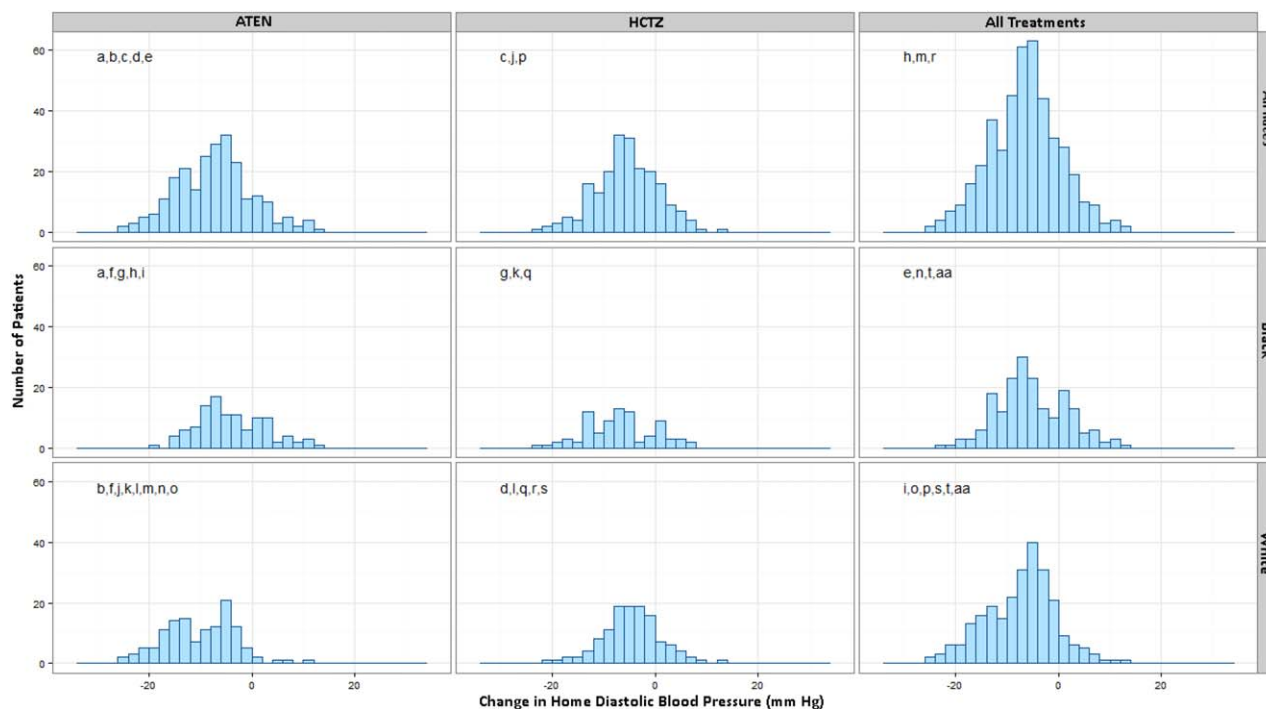


Figure 1 Histograms of change in HDBP in participants treated with HCTZ or ATEN. Each panel represents the distribution of change in HDBP with columns stratifying groups by treatment of ATEN, HCTZ, or combining all participants. Rows combine all participants or stratify by race. Panels with matching letters represent groups that displayed statistically significant differences in HDBP response ($q < 0.1$).

The eight most significantly changed metabolites in the white group treated with HCTZ were uric acid, ribonic acid, 1-hexadecanol, kynurenine, glycerol-gulo-heptose NIST, dihydroabietic acid, behenic acid, and glucose-1-phosphate (**Table 1**). Four metabolites were significantly increased ($q < 0.1$) in the black group treated with HCTZ (uric acid, propane-1-2-3-tricarboxylate NIST, 223865, 455836). Although four metabolites were significantly decreased ($q < 0.1$) in the white group treated with HCTZ, glycine was the only identified metabolite (glycine, 217797, 294266, 228983). Overall, there were five metabolites significantly decreased ($q < 0.1$) in the black group treated with HCTZ composed of phytol, 428330, 200906, 617225, and phosphoethanolamine (**Table 1**).

Clustering of significant metabolites in signature of exposure

Twenty-nine unique metabolites at baseline were significantly associated with ATEN exposure in either the black, white, or both groups ($q < 0.2$) and were clustered together using Modulated Modularity Clustering.²⁰ A group of eight metabolites (300379, 240264, nicotinic acid, 239995, 210904, 566053, 210901, and 516629) were clustered together with an absolute average correlation of 0.43. The remaining cluster, composed of the additional 21 metabolites, had an average absolute correlation of 0.11 (**Supplementary Figure S1**).

A total of 170 unique metabolites were associated at baseline with HCTZ treatment in the white, black, or both groups ($q < 0.1$). Overall, 5 clusters were identified with 10, 6, 5, 8, and 141 metabolites, respectively, and absolute

average correlation coefficients of 0.78, 0.77, 0.73, 0.65, and 0.11, respectively (**Figure 2**).

Signature of response to ATEN and HCTZ Baseline metabolite associations with HDBP response.

Patient distributions of HDBP response confirmed that, although most participants in each group experienced lower HDBP after treatments with ATEN or HCTZ, there was a statistically significant difference between the responses observed in black and white groups for either treatments (**Figure 1**). Initial tests were conducted to determine the associations of baseline metabolites with changes in HDBP. The 5-methoxytryptamin was the only metabolite at baseline that was associated with a change in HDBP for the white group treated with ATEN ($q < 0.2$), and exhibited a negative association. However, seven metabolites were associated at baseline with a change in HDBP for the black group treated with ATEN, with all but three being unidentified (2,4-diaminobutyric acid, arabitol, and O-acetylserine; **Table 2**).

Arachidonic acid and 223548 were the only metabolites associated at baseline with change in HDBP for the white group treated with HCTZ ($q < 0.2$), and both displayed positive associations. An unidentified metabolite, 223548, was also significantly associated with a change in HDBP at baseline in the black group treated with HCTZ. However, unlike in the white cohort, 223548 was negatively associated with change in HDBP in the black cohort (**Table 2**).

Metabolite changes associated with HDBP response. Metabolite change from baseline to posttreatment was tested

Table 1 Hydrochlorothiazide: metabolic signature of exposure ($q < 0.2$)

Direction of association	White participants ($n = 123$)					Black participants ($n = 83$)						
	Metabolite	Pretreatment median	Posttreatment median	Difference	P value	q value	Metabolite	Pretreatment median	Posttreatment median	Difference	P value	q value
Increased^b	Uric acid	50,143	62,808	12,665	3.93E-11	1.91E-08	Uric acid	48,159	62,892	14,733	1.83E-08	8.90E-06
	Ribonic acid	6.33 ^a	6.48 ^a	0.15 ^a	7.48E-07	0.0002	Propane-1,2,3-tricarboxylate NIST	6.73 ^a	7.05 ^a	0.32 ^a	0.0006	0.0472
	228,583	6.17 ^a	6.39 ^a	0.22 ^a	3.52E-06	0.0006	223,865	6.52 ^a	6.84 ^a	0.32 ^a	0.0006	0.0472
	338,896	566	670	104	8.80E-06	0.0011	455,836	792	928.50	136.5	0.0010	0.0612
	1-Hexadecanol	6.16 ^a	6.31 ^a	0.15 ^a	1.77E-05	0.0014	470,983	7.85 ^a	8.52 ^a	0.67 ^a	0.0038	0.1682
	Erythritol	2,900	3,224	324	1.73E-05	0.0014	571,392	7.31 ^a	7.48 ^a	0.17 ^a	0.0044	0.1780
	Kynurenine	6.31 ^a	6.42 ^a	0.10 ^a	2.54E-05	0.0018	340,252	1,087	1,393	306	0.0048	0.1815
	Glycero- gulo-heptose NIST	6.37 ^a	6.52 ^a	0.15 ^a	2.93E-05	0.0018	339,431	636	821	185	0.0053	0.1860
	223,597	380.5	450	69.5	4.89E-05	0.0026						
	223865	6.38 ^a	6.79 ^a	0.42 ^a	6.56E-05	0.0032						
	Dihydrobiotic acid	6.76 ^a	6.85 ^a	0.09 ^a	7.33E-05	0.0032						
	2-ketoisocaproic acid major	1,061	1,378	317	9.97E-05	0.0040						
	Aconitic acid	385	480.50	95.50	0.0001	0.0040						
	Behenic acid	1,437	1,583	146	0.0001	0.0043						
	Glucose 1-phosphate	6.37 ^a	6.45 ^a	0.085 ^a	0.0001	0.0043						
	Glycine	12.08 ^a	12 ^a	-0.08 ^a	0.0066	0.0340						
	217,797	579	565	-14	0.0072	0.0358						
	294,266	6.85 ^a	6.79 ^a	-0.06 ^a	0.0149	0.0600						
	228,983	6.23 ^a	6.17 ^a	-0.06 ^a	0.0277	0.0922						
	428,330	7.75 ^a	7.70 ^a	-0.05 ^a	0.0491	0.1344	Phytol	6.74 ^a	6.59 ^a	-0.15 ^a	2.98E-05	0.0073
Threonine	47,018	43,279	-3,739	0.0521	0.1394	428,330	7.82 ^a	7.73 ^a	-0.09 ^a	0.0003	0.0460	
Aminomalonic acid	8.31 ^a	8.25 ^a	-0.06 ^a	0.0552	0.1443	200,906	869	758	-111	0.0004	0.0472	
218,951	6.42 ^a	6.36 ^a	-0.06 ^a	0.0556	0.1443	617,225	6.78 ^a	6.68 ^a	-0.10 ^a	0.0007	0.0472	
446,067	30,594	27,414	-3,180	0.0585	0.1484	Phosphoethanolamine	7.34 ^a	7.18 ^a	-0.16 ^a	0.0018	0.0945	
Glutamine	201,749	188,413	-13,336	0.0686	0.1662	Methoxytyrosine NIST	6.43 ^a	6.22 ^a	-0.21 ^a	0.0028	0.1356	
226,846	6.96 ^a	6.85 ^a	-0.11 ^a	0.0729	0.1724							
Serine	39,561	38,059	-1,502	0.0798	0.1834							

^aValues for these metabolites are log-transformed.

^bTop 15 most significant metabolites presented here of a total of 201 metabolites ($q < 0.2$).

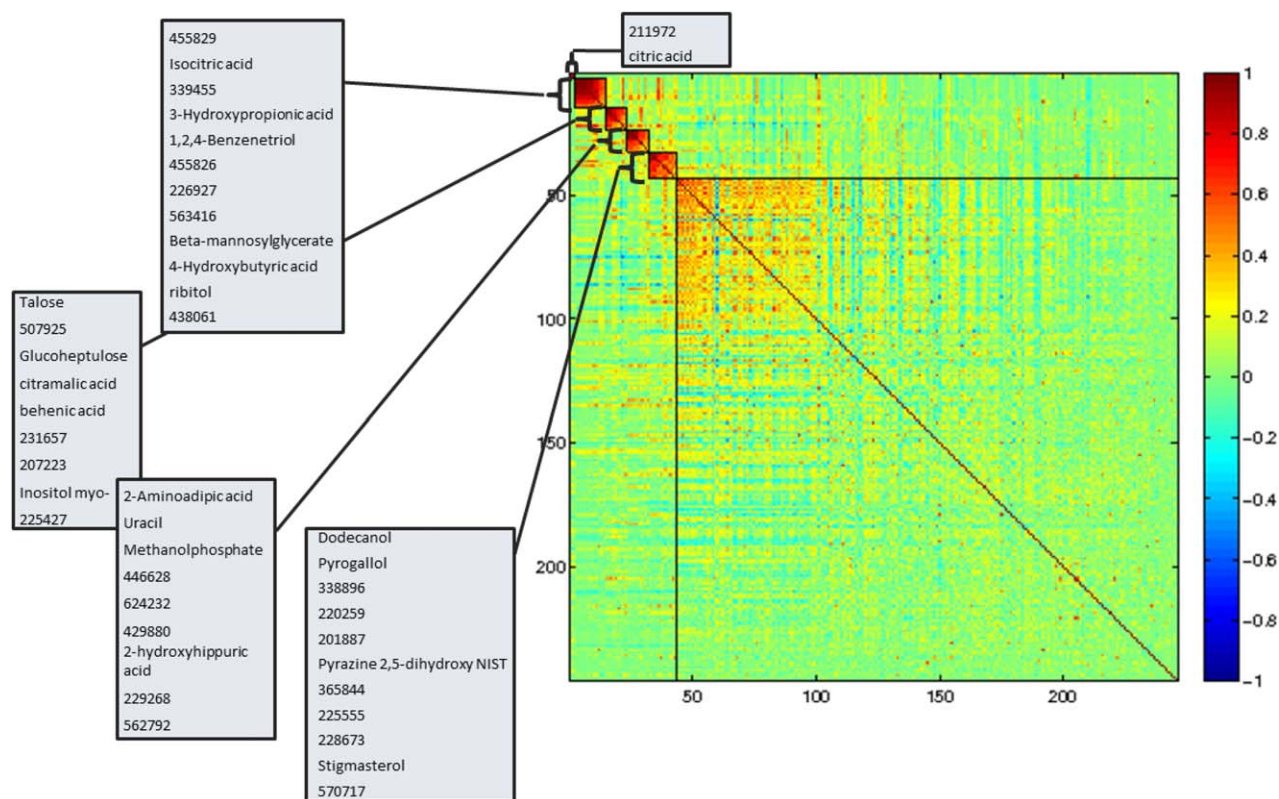


Figure 2 Hierarchical clustering of metabolites altered upon treatment of HCTZ ($q < 0.2$). The boxes on the heat map outline significant modules and are annotated with the metabolites within each module. The x-axis and y-axis are metabolites and the shade of color represents the Pearson correlation.

for associations with HDBP response. In the white group treated with ATEN, changes in 4 metabolites (2 known: 2-oxogluconic acid NIST and maltose) were significantly associated with a HDBP change ($q < 0.05$). In the black group treated with ATEN, the change in six metabolites was associated with HDBP response with four identified metabolites (isothreonic acid, gluconic acid, 4-hydroxyproline, and indole-3-acetate; **Table 3**).

Participants in the white group treated with HCTZ displayed no metabolite changes that were significantly associated with HDBP response at $q < 0.05$. In the black group treated with HCTZ, there was only a single, unidentified metabolic change (437822) that was associated with an HDBP response ($q < 0.05$; **Table 3**).

Multivariable model predictive of HDBP response.

Using baseline metabolite profiles, models were developed to predict HDBP response for participants treated with either ATEN (**Supplementary Figure S2**) or HCTZ (**Supplementary Figure S3**) in order to identify metabolites that are associated with a good BP response regardless of the mode of therapy used. Models combining groups by either race or drug treatment performed better, presumably from increased statistical power with the increased number of participants. Models for All treatments-All races, ATEN-All races, HCTZ-All races, and All treatments-White were statistically significant for both discovery and validation sets with a multiple test corrected $P < 0.006$ with R^2 values for

the validation set of 0.19, 0.43, 0.49, and 0.33, respectively (**Supplementary Figures S2, S3, and Figure 3**). A summary of all results for the discovery and validation data can be seen in **Figure 3**.

Additional multivariable models were constructed using changing metabolites and are described in the supplementary material (**Supplemental Figures S4–S6**).

Pathway analysis

Pathways enriched by metabolite signature of drug exposure.

Pathway analysis detected 11 significant pathways for both ATEN and HCTZ (adjusted $P < 0.05$). In subjects treated with ATEN, pathway analysis detected significant decreases of the α -linolenic acid and linoleic acid metabolism pathway (adjusted $P = 1.09 \times 10^{-8}$ for all races and 7.57×10^{-5} for whites; **Table 4**). In addition, the fatty acid biosynthesis (adjusted $P = 0.0055$) and glycerolipid metabolism (adjusted $P = 0.0005$) pathways were significantly decreased when both races were combined, and when only the white group was tested (i.e., adjusted $P = 0.0093$ and $P = 0.0023$, respectively).

Interestingly, all significant pathways in the HCTZ treatment group were due to increases in metabolite levels. The purine metabolism pathway was significantly increased for all races (adjusted $P < 1 \times 10^{-9}$), the white group (adjusted $P = 2.24 \times 10^{-8}$), and the black group (adjusted $P = 5.12 \times 10^{-5}$). This was the only pathway significantly affected when the data was

Table 2 Baseline metabolites associated with change in HDBP ($q < .20$)

Race	Atenolol				Hydrochlorothiazide			
	Metabolite	P value	q value	Association	Metabolite	P value	q value	Association
All	O-acetylserine ^a	0.001	0.030	Negative	Palmitoleic acid ^a	0.003	0.086	Positive
	483390 ^a	0.001	0.041	Positive	Salicylic acid ^a	0.007	0.007	Positive
	213253 ^a	0.003	0.081	Positive				
	6-deoxyglucitol NIST ^a	0.008	0.167	Positive				
	Beta-mannosylglycerate ^a	0.009	0.177	Positive				
	Fucose + rhamnose ^a	0.009	0.180	Positive				
	216043 ^a	0.011	0.198	Positive				
White	5-Methoxytryptamine ^b	0.009	0.176	Negative	Arachidonic acid ^b	0.002	0.067	Positive
					223548 ^b	0.007	0.153	Positive
Black	2,4-diaminobutyric acid ^c	0.002	0.068	Negative	223548 ^c	0.004	0.11	Negative
	470983 ^c	0.002	0.074	Negative				
	340257 ^c	0.003	0.078	Negative				
	228377 ^c	0.003	0.094	Positive				
	Arabitol ^c	0.004	0.101	Positive				
	O-acetylserine ^c	0.004	0.105	Negative				
	483390 ^c	0.004	0.107	Positive				

HDBP, home diastolic blood pressure.

^aAdjusted for: gender, baseline glucose levels, baseline HDBP.^bAdjusted for: baseline HDBP and baseline renin.^cAdjusted for: baseline renin and baseline glucose levels.

stratified by only black subjects. For HCTZ exposure, other significantly increased pathways include galactose metabolism, lactose synthesis, gluconeogenesis, glycolysis, and the urea cycle when both races were combined (adjusted $P < 0.05$). When stratified by only white subjects, significant increases in lactose synthesis, plasminogen synthesis, glycolysis, tryptophan metabolism, galactose metabolism, and gluconeogenesis were observed with adjusted $P < 0.05$ for each pathway.

DISCUSSION

The results of the metabolomics analysis in this study reveal important insights into interindividual and interpopulation differences in response to antihypertensives. HCTZ and ATEN have distinct mechanisms of action. HCTZ, a thiazide diuretic, lowers BP by inhibiting renal reabsorption. ATEN, on the other hand, is a beta-blocker. The

Table 3 Metabolite changes associated with change in HDBP

Race	Atenolol				Hydrochlorothiazide			
	Metabolite	P value	q value	Association	Metabolite	P value	q value	Association
All	Indole-3-acetate ^a	4.47E-06	0.0005	Negative	Palmitoleic acid ^d	0.003	0.086	Positive
	Gluconic acid ^a	7.23E-05	0.005	Negative	Salicylic acid ^d	0.007	0.007	Positive
	Inositol myo- ^a	8.75E-05	0.006	Negative				
	339186 ^a	0.0003	0.017	Negative				
	226927 ^a	0.0008	0.033	Negative				
	2-oxogluconic acid NIST ^a	0.0009	0.037	Positive				
	199773 ^a	0.001	0.043	Negative				
White	2-oxogluconic acid NIST ^b	1.89E-06	0.0002	Positive				
	Inositol myo- ^b	0.0001	0.008	Negative				
	270066 ^b	0.0002	0.010	Positive				
Black	Maltose ^b	0.0004	0.020	Negative				
	Erythritol ^c	0.0001	0.007	Negative	437822 ^e	0.003	0.028	Negative
	Isothreonic acid ^c	0.0001	0.009	Negative				
	Gluconic acid ^c	0.0002	0.012	Negative				
	228377 ^c	0.0002	0.012	Positive				
	4-hydroxyproline ^c	0.0003	0.017	Positive				
Indole-3-acetate ^c	0.001	0.046	Negative					

HDBP, home diastolic blood pressure; HOMA, homeostasis model of assessment.

^aAdjusted for: race, baseline renin, baseline HOMA, baseline triglycerides.^bAdjusted for: baseline HDBP and baseline renin.^cAdjusted for: baseline renin and baseline glucose levels.^dAdjusted for: gender, baseline HDBP, baseline glucose levels.^eAdjusted for: gender and baseline glucose levels.

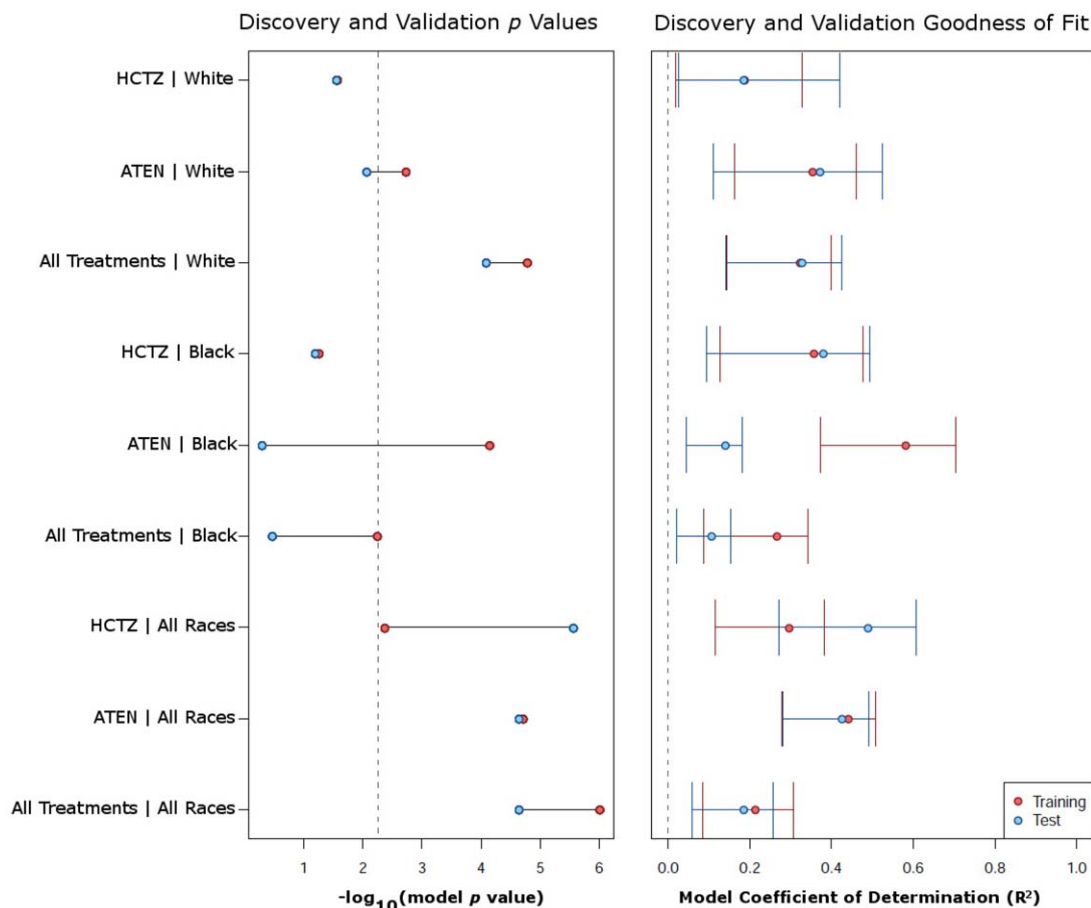


Figure 3 Multivariable modeling of change in HDBP using baseline metabolic signatures. (a) Shows for each model the p value obtained for the discovery (red) and validation (blue) sets. The vertical dashed line represents the Bonferroni corrected P value required for the model results to be considered statistically significant. The x-axis is the $-\log_{10}(P \text{ value})$ so that points to the right represent increasing significance. (b) The R^2 of each model separated by discovery (red) and validation (blue) sets. The error bars represent the bootstrapped 95% confidence interval.

mechanism by which ATEN lowers BP has not been fully elucidated, although it is thought that the primary mechanism is through reduced cardiac output via a reduction in heart rate. The mechanistic differences in these drugs are reflected in the distinctive metabolite signatures detected here. Furthermore, the observed heterogeneity of response between white and black groups treated with either ATEN or HCTZ, based on HDBP ($q < 0.05$), stresses the need for an improved understanding of the mechanisms of action underlying these two medications (Figure 1).

Treatment with HCTZ resulted in a substantially different metabolic profile compared to treatment with ATEN (Table 1 and Supplementary Table S1). Uric acid was significantly increased upon treatment with HCTZ in both black and white groups consistent with the well-documented side effect of hyperuricemia of thiazide-based diuretics.^{23,24} This metabolomic finding confirms the previous clinical findings of uric acid increase in this same cohort,²⁵ but expand on the previous findings with a more comprehensive interrogation of the “metabolome.” The effect observed on uric acid may be indicative of the renal mechanism of action of HCTZ, which is not observed with ATEN treatment.

Several baseline or changing metabolites demonstrated significant positive or negative associations with HDBP response in participants treated with ATEN or HCTZ. Both baseline levels and the change of palmitoleic acid were positively associated with an HDBP response in participants treated with HCTZ, when all races were combined. Although the role of palmitoleic acid in BP is not well understood, several epidemiological studies have observed significant associations relating increases in palmitoleic acid with increased BP.^{26,27} In white participants treated with ATEN, baseline 5-methoxytryptamine displayed a negative association with HDBP response. In black participants treated with ATEN, change in indole-3-acetate was associated with HDBP response. The 5-methoxytryptamine is a tryptamine derivative related to serotonin. Also, melatonin and indole-3-acetate are derived from tryptophan and are important components of the gut microbiome in humans.^{28,29} These two metabolites suggest a possible impact of tryptophan or tryptamines (e.g., serotonin, melatonin) on antihypertensive response to multiple classes of antihypertensive drugs (e.g., beta-blockers and thiazide diuretics), and a possible impact for the gut microbiome.³⁰ Additional studies have identified metabolites

Table 4 Results from correlated Lancaster pathway analysis signature of exposure

Pathway name	Number of metabolites in:					P value ^f
	SMPDB ID ^a	KEGG ID ^b	Drug	Race	Direction of change	
Purine metabolism	SMP00050	map00230	HCTZ	All	Increase	<1 × 10 ⁻⁹
Alpha linolenic acid and linoleic acid metabolism	SMP00018	map00592	ATEN	All	Decrease	1.09 × 10 ⁻⁰⁸
Purine metabolism	SMP00050	map00230	HCTZ	White	Increase	2.24 × 10 ⁻⁰⁶
Alpha linolenic acid and linoleic acid metabolism	SMP00018	map00592	ATEN	White	Decrease	7.57 × 10 ⁻⁰⁷
Purine metabolism	SMP00050	map00230	HCTZ	Black	Increase	5.12 × 10 ⁻⁰⁵
Galactose metabolism	SMP00043	map00052	HCTZ	All	Increase	0.0003
Gluconeogenesis	SMP00128	map00010	HCTZ	All	Increase	0.0004
Glycerolipid metabolism	SMP00039	map00561	ATEN	All	Decrease	0.0005
Glycolysis	SMP00040	map00010	HCTZ	All	Increase	0.0009
Glycolysis	SMP00040	map00010	HCTZ	White	Increase	0.0013
Gluconeogenesis	SMP00128	map00010	HCTZ	White	Increase	0.0023
Glycerolipid metabolism	SMP00039	map00561	ATEN	White	Decrease	0.0025
Fatty acid biosynthesis	SMP00456	map00052	ATEN	All	Decrease	0.0055
Lactose synthesis	SMP00444	map00010	HCTZ	All	Increase	0.0065
Fatty acid biosynthesis	SMP00456	map00052	ATEN	White	Decrease	0.0093
Plasmalogen synthesis	SMP00479	map00038	HCTZ	White	Increase	0.0101
Tryptophan metabolism	SMP00063	map00052	HCTZ	White	Increase	0.0133
Galactose metabolism	SMP00043	map00052	HCTZ	White	Increase	0.0146
Lactose synthesis	SMP00444	map00010	HCTZ	White	Increase	0.0258
Urea cycle	SMP00059	map00330	HCTZ	All	Increase	0.0377

^aSmall Molecule Pathway Database ID.^bKyoto Encyclopedia of Genes and Genomes ID.^cNumber of metabolites in pathway.^dNumber of metabolites studied.^eNumber of metabolites studied that also overlapped with metabolites in the pathway.^fBonferroni corrected *P* value.

associated with BP that point toward potential mechanisms in the gut microbiome.¹³ These results are further supported by the significant effect on the tryptophan metabolism pathway in white participants treated with HCTZ (adjusted $P < 0.05$; **Table 4**). Furthermore, several studies have demonstrated the ability for some tryptamines to impact BP and their role as either vasopressors or vasodepressors.^{31–33}

Treatment with either ATEN or HCTZ resulted in several significantly altered pathways. Namely, effects on fatty acid biosynthesis and glycerolipid metabolism were observed in subjects treated with ATEN (**Table 4**). Effects on purine metabolism (driven mainly by effects on uric acid), galactose metabolism, lactose synthesis, plasmalogen synthesis, glycolysis, gluconeogenesis, and tryptophan metabolism were observed in subjects treated with HCTZ (adjusted $P < 0.05$; **Table 4**). Plasmalogens, a type of phospholipid, play an important role in cell signaling, membrane structure, and are protective against reactive oxygen species.^{34,35} Increases in plasmalogens have been considered markers of increased oxidative stress and have been associated with increased cardiovascular mortality in participants with endstage renal disease.³⁴ Several metabolites implicated in fatty acid biosynthesis, glycerolipid metabolism, and alpha linolenic acid and linoleic acid metabolism clustered after ATEN exposure (**Supplementary Figure S1**). Many unknown metabolites were highly clustered after HCTZ exposure (**Figure 2**). Annotating these unknown metabolites will be very important and may provide new insight into biological pathways impacted by HCTZ treatment.

For many years, studies have noted glucose impairment in some patients treated with thiazide-based diuretics.^{36,37} Although still under debate, recent studies have also demonstrated associations with thiazide-based treatments and increased risk of type II diabetes and subsequently increased the risk of cardiovascular events.^{38,39} Phosphoethanolamines are used in the construction of sphingolipids, and perturbations in sphingolipid metabolism have been implicated in adverse cardiovascular effects, have demonstrated the ability to alter insulin resistance, and in turn may be connected to increased risk of type II diabetes.^{40–42} We detected a statistically significant decrease in circulating O-phosphoethanolamine in the black group exposed to HCTZ ($q < .05$), which may be due to an increase in sphingolipid biosynthesis. Lipidomic profiling, with increased coverage of the sphingolipid metabolism pathway, will be required to further elucidate the impact of HCTZ treatment on this pathway.

Additionally, increases in metabolic sugars resulted in increases in galactose metabolism, lactose synthesis, glycolysis, and gluconeogenesis pathways (**Table 4**). This increase in metabolic sugars may also exacerbate risks of glucose impairment in patients treated with HCTZ. However, because of the nontargeted approach of this metabolite panel, many of these metabolites in our panel had only minimal overlap with any specific pathway. Therefore, a more targeted approach, with improved coverage of particular pathways, is needed to follow-up these new hypotheses and gain a better understanding of how these biological pathways are related to treatment response.

Multivariable modeling offers the advantage of exploring combinations of factors that may provide predictive capability

for patient drug response. Multivariable modeling in the current study highlighted a subset of baseline metabolic measurements that were predictive for ATEN (validation set: $P = 2.27 \times 10^{-5}$, $R^2 = 0.43$) and HCTZ (validation set: $P = 2.74 \times 10^{-6}$, $R^2 = 0.49$) induced HDBP response when both black and white groups were combined (**Supplementary Figures S2–S3, Figure 3**). The validation set was tested using the model constructed on the discovery data, in order to assess model overfitting, and models that were statistically significant in both the discovery and the validation set would be expected to have strong predictive performance. Models were not statistically significant when treatments and race were separated, presumably from the reduction in sample size and subsequent loss of statistical power. Race was a significant covariate that was retained in the model for ATEN during feature selection; however, gender was a significant covariate included in the model for HCTZ treatment. Together, these models demonstrate that a significant contribution of the change in HDBP by ATEN and HCTZ can be explained by metabolomics profiling. However, many of the metabolites included as features in the models are currently unidentified, underscoring the need to identify and characterize these metabolites. With extensive additional validation, these models could provide a cost-effective, non-invasive, diagnostic capable of determining which treatment is the most likely to elicit a positive outcome for a patient.

In addition to being significantly associated with HDBP in white participants treated with ATEN, baseline methoxytryptamine was also selected as a variable in the multivariable model for ATEN for white participants and when both black and white participants were combined. Baseline levels of palmitoleic acid and oleic acid were both included as features for predicting HCTZ-induced changes in HDBP. The coefficient for palmitoleic acid was positively associated with change in HDBP; whereas, the coefficient for oleic acid, albeit small, was negatively associated with change in HDBP, consistent with previous studies on the effects of these fatty acids on BP.^{26,43}

Through the use of metabolomics, we have identified sets of metabolites significantly altered by either treatment with HCTZ or ATEN and are associated with change in HDBP. In addition, we have demonstrated that racial differences in response to either drug exist and can be, at least partially, explained through metabolic profiles. Although additional work will be needed to determine the full impact of these medications on participants, these data offer new insights into the impact of hypertensive treatments on fatty acid and other metabolic pathways related to BP response.

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R.C.D., R.F.F., Y.G., and A.M.R. designed the research. D.M.R., A.L.B., R.C.D., M.S., R.F.F., L.W., H.Z., and Y.G. performed the research. D.M.R., M.M., S.B.G., M.S., L.W., and H.Z., analyzed the data. M.E.S. and O.F. contributed new reagents/analytical tools.

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