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Metabolic phenotyping for discovery of urinary biomarkers of diet, xenobiotics and blood pressure in the INTERMAP Study: An overview

Queenie Chan^{1,2}, Ruey Leng Loo^{3,4}, Timothy M. D. Ebbels^{2,4}, Linda Van Horn⁵, Martha L. Daviglius^{5,6}, Jeremiah Stamler⁵, Jeremy K. Nicholson^{4,7}, Elaine Holmes^{2,4,7}, Paul Elliott^{1,2,7}

¹Department of Epidemiology and Biostatistics, School of Public Health, Imperial College London, London, UK

²MRC-PHE Centre for Environment and Health, School of Public Health, Imperial College London, London, UK

³Medway Metabonomics Research Group, Medway School of Pharmacy, Universities of Kent and Greenwich, Chatham Maritime, Kent, UK

⁴Division of Computational and Systems Medicine, Department of Surgery and Cancer, Imperial College London, London, UK

⁵Department of Preventive Medicine, Feinberg School of Medicine, Northwestern University, Chicago, IL, USA

⁶Institute for Minority Health Research, University of Illinois at Chicago, Chicago, IL, USA

⁷MRC-NIHR National Phenome Centre, Imperial College London, London, UK

Corresponding author: Queenie Chan or Paul Elliott

Department of Epidemiology and Biostatistics, School of Public Health, Imperial College London, Norfolk Place, London, W2 1PG, UK.

Tel: +44 207 594 3328. Fax: +44 207 594 3456.

Email: g.chan@imperial.ac.uk or p.elliott@imperial.ac.uk

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Abstract

The aetiopathogenesis of cardiovascular diseases (CVD) is multifactorial. Adverse blood pressure (BP) is a major independent risk factor for epidemic CVD affecting about 40% of the adult population worldwide and resulting in significant morbidity and mortality. Metabolic phenotyping of biological fluids has proven its application in characterising low molecule weight metabolites providing novel insights into gene-environmental-gut microbiome interaction in relations to a disease state. In this review, we synthesise key results from the International Study of Macro/Micronutrients and Blood Pressure (INTERMAP) Study, a cross-sectional epidemiological study of 4,680 men and women aged 40-59 years from Japan, the People's Republic of China, the United Kingdom, and the United States. We describe the advancements we have made on: 1) analytical techniques for high throughput metabolic phenotyping; 2) statistical analyses for biomarker identification; 3) discovery of unique food-specific biomarkers; and 4) application of metabolome-wide association (MWA) studies to gain a better understanding into the molecular mechanisms of cross cultural and regional BP differences.

Introduction

Adverse blood pressure (BP) (prehypertension and hypertension) is a major independent risk factor for epidemic cardiovascular diseases (CVD), affecting about 40% of the adult population worldwide.¹⁻⁴ Its causation is multifactorial encompassing both environmental -- diet, other aspects of lifestyle -- and genetic factors. Public health policies aiming to improve the prevention of high BP and/or maintain optimal BP level typically involve efforts to tackle known modifiable risk factors such as reduction of high salt intake, moderation of alcohol intake, maintenance of normal weight and increased physical activity. While large-scale genome-wide association studies (GWAS) have identified common variants, novel loci and pathways associated with BP,⁵⁻⁹ the genetic contribution to BP variation in the population is modest.^{5, 7, 8} Despite the effectiveness of non-pharmacological approaches to BP control, many people with high BP are reliant on antihypertensive drugs or BP remains untreated or poorly controlled.^{10, 11}

The INTERnational study of MACro/micronutrients and blood Pressure (INTERMAP) is a cross-sectional epidemiological study of 4,680 men and women aged 40-59 years from Japan, the People's Republic of China, the United Kingdom, and the United States (www.clinicaltrials.gov NCT00005271).¹² The main aim of the INTERMAP Study is to investigate the aetiology of adverse BP with emphasis on studying diet-BP associations (**Figure 1**).^{12, 13} Participants were selected randomly from population lists, stratified by age/sex. Each participant attended four visits, visits 1 and 2 on consecutive days, visits 3 and 4 on consecutive days on average 3 weeks later. At each of the four visits, BP was measured twice with a random zero sphygmomanometer and dietary data were collected by a trained certified interviewer with use of the in-depth multi-pass 24-h recall method.¹⁴ Height and weight were measured twice on the first and third visits. On the first visit, questionnaire data were obtained on demographic, medical history, medication intake, and other possible confounders. Each participant provided two 24-h urine collections, start and end timed at the research centre. The two timed 24-hour urine collections per person on each of the 4,680 INTERMAP participants are a pivotal feature of study design that enabled the introduction of metabolome-wide association (MWA) study (see later section). The study received institutional ethics committee approval for each site; all participants gave written informed consent.

In the past decade, the INTERMAP Study, has added new knowledge to the limited data previously available on the effects of nutritional factors on BP.¹⁵ These advances include the inverse relationship between BP and intakes of vegetable protein,¹⁶ glutamic acid,¹⁷ total and insoluble fibre,¹⁸ raw and cooked vegetables,¹⁹ total polyunsaturated fatty acids (PFA) and linoleic acid,²⁰ oleic acid from vegetable sources,²¹ total omega-3 fatty acids and linolenic acid,²² phosphorus (P), calcium (Ca) and magnesium (Mg),²³ non-heme iron (Fe) and total Fe,²⁴ and starch²⁵; and the direct associations of sugars (fructose, glucose and sucrose),²⁶ cholesterol,²⁷ glycine and alanine,²⁸ raw fruits,²⁹ and oleic acid from animal sources.²¹ The INTERMAP Study showed that diet-induced metabolic acidosis was positively associated with BP (not significant after controlling for body mass index, BMI);³⁰ whereas a cohort of over 61,000 persons reported a positive association of metabolic acid load (such as serum bicarbonate, urine acidity) and cardiovascular mortality.³¹ While the INTERMAP Study reported a small non-significant inverse relationship between urinary Mg and BP³²; the World Health Organization-coordinated Cardiovascular Diseases and Alimentary Comparison (CARDIAC) Study showed urinary Mg to creatinine ratio was inversely associated with CVD risk factors such as BMI and BP.³³ Although important advances have been made by the INTERMAP Study in improving understanding of the aetiology of high BP, together with other research into the physiology of BP regulation such as the control of kidney fluid and salt balance *via* the renin-angiotensin-aldosterone system,³⁴⁻³⁶ sympathetic nervous system activity,^{37, 38} and the role of structure and function of blood vessels,³⁹ gaps still remain in our knowledge of the causes and mechanisms of adverse BP levels. New approaches are needed to enhance understanding of the multifactorial aetiopathogenesis of BP.

High resolution proton nuclear magnetic resonance (NMR) spectroscopy have been successfully applied for the investigation of drug metabolism and toxicology as well as disease development within a biological system, using biological fluids such as urine, plasma/serum as well as bile, cerebrospinal fluids and dialysates.^{40, 41} Recent studies have shown the importance of the gut-microbiome in the aetiology of a number of chronic diseases such as atherosclerosis, diabetes and the metabolic syndrome, obesity as well as raised BP.⁴²⁻⁴⁷ Metabolic phenotyping of biological fluids using spectroscopic methods enables the investigation of gene-environmental-gut microbiome interactions on disease risk and is therefore an attractive approach for gaining new insights into BP mechanisms and its

associated pathways.^{45, 48-50} The INTERMAP Study has capitalised on the evolving technologies in metabolic phenotyping to enhance its rich nutrient and anthropometric data by incorporating this top down system approach to investigate the association of BP and urinary markers that are linked to environmental exposures including diets and xenometabolomes.⁴⁸⁻⁵² This review demonstrates the key progress made by the INTERMAP Study and the introduction of the metabolome-wide association (MWA) study on diet and BP.¹³ However, readers may wish to refer to other recent reviews on the principles of metabolic phenotyping.⁵³⁻⁵⁵

Analytical method development

Metabolic phenotyping involves the application of high-throughput advanced spectroscopic methods, such as proton nuclear magnetic resonance (¹H NMR) spectroscopy and mass spectroscopy (MS), to biological samples. In INTERMAP comprehensive metabolic phenotyping by ¹H NMR spectroscopy of the two 24-h urine samples from each participant (N=4,630) has been performed.¹³ The repeatability, accuracy and stability of the ¹H NMR spectral profiles were evaluated and overall analytical reproducibility of the ¹H NMR was found to be >98%.⁵⁶ In INTERMAP, boric acid (borate, a preservative known to bind covalently to vicinal diols and some amino acids) was added to the urine collection jars in the field to prevent bacterial overgrowth. The effect of the boric acid on the ¹H NMR urinary spectra was also assessed.^{37, 38} It was shown that the overall changes in the urinary ¹H NMR spectral profile caused by borate addition were negligible compared with the physiological and metabolic differences between individuals.⁵⁷ These studies have led to recommendations for improved sample preparation and handling of urine samples, as well as processing methods for large-scale epidemiologic research.⁵⁸

In addition to metabolic phenotyping via ¹H NMR spectroscopy, we obtained extensive data on 20 urinary amino acids using conventional cation-exchange chromatography followed by post-column derivatisation (Biochrom 20 and Biochrom 30). These data provide a unique population based dataset on urinary amino acid excretion levels in different populations. We then applied gas chromatography mass spectrometry (GC-MS) and liquid chromatography-tandem mass spectrometry (LC-MS/MS) for high throughput urinary amino acid analysis and compared their sample preparation, run-time, number of analytes amenable to

quantification, cost, limit of quantification (LOQ), reproducibility and validity with conventional amino acid analysis.⁵⁹ The amount of urine needed for GS-MS and LC-MS/MS were 40-50 μ L, much less than the 200 μ L required for the amino acid analyser. Moreover, the run-time for the amino acid analyser was approximately 5 to 6 times longer than that of GC-MS and LC-MS/MS. The Pearson correlation coefficients of amino acids measured by GC-MS and the amino acid analyser ranged from 0.80 (tryptophan) to 0.98 (glycine); correlation coefficients comparing LC-MS/MS with the amino acids analyser ranged from 0.56 (arginine) to 0.95 (lysine). Our findings showed that GC-MS offered higher reproducibility and completely automated sample pre-treatment compared with LC-MS/MS and conventional amino acid analysis, which on the other hand covered more amino acids and related amines.

We also applied ultra performance liquid chromatography triple quadruple tandem mass spectrometry (UPLC-TQ-MS/MS) for simultaneous detection (both positive and negative electrospray ionization modes) and quantification of three gut microbial co-metabolites, phenylacetylglutamine (PAG), 4-cresyl sulphate and hippurate in the urinary specimens from 2,000 INTERMAP US participants.⁶⁰ This targeted high-throughput UPLC-TQ-MS/MS method was developed specifically to measure these gut microbial co-metabolites, which may be implicated in obesity⁶¹ and other chronic diseases such as cardiovascular and kidney disease.^{62,63} Following the US Food and Drug Administration (FDA) guidelines, the imprecision (CV: coefficient of variation) and inaccuracy (recovery) of the method were assessed using replicates of quality control (QC) urine samples at different concentration levels. The CV and recovery were found to be within acceptable limits of FDA guidelines. The study demonstrated the applicability of metabolic phenotyping by UPLC-TQ-MS/MS in a large scale epidemiological study.

Statistical method development

Metabolomic data pose special challenges to statistical analysis, including high dimensionality, strong co-linearity, non-linear and highly complex spectral profiles, and presence of structured and unstructured noise (due to within- and between-individual variability).⁶⁴⁻⁶⁸ Spectroscopic data are first pre-processed, including spectral alignment and normalisation of the full-resolution spectral data. Multiple statistical strategies are then applied, including use of both unsupervised and supervised multivariate data analysis

techniques to extract information from the data.⁶⁹ Principal component analysis (PCA) is routinely used to identify main sources of variation in the data, and detect outlying values with the goal of providing the most compact representation of the data.^{64, 67-69} Hierarchical cluster analysis (HCA) is another method widely used in exploratory data analysis to provide an overview of the data by grouping phenotypes according to their similarity, without assuming any prior knowledge of the data.^{64, 68, 69} Other techniques include partial least squares (PLS), orthogonal partial least squares (O-PLS), and orthogonal partial least squares discriminant analysis (OPLS-DA)⁷⁰⁻⁷² aiming to extract discriminatory metabolic signals from the datasets are often applied after the initial exploratory analysis; whilst statistical spectroscopy methods, accommodating complex spectral data structure and correlations, such as statistical total correlation spectroscopy (STOCSY),^{73, 74} iterative-STOCSY (I-STOCSY),⁷⁵ cluster analysis statistical spectroscopy (CLASSY),⁷⁶ and subset optimization by reference matching (STORM),⁷⁷ are used to aid the biomarker identification process. Statistical Heterospectroscopy (SHY) is a statistical strategy for the analysis of multiple spectroscopic datasets e.g. ¹H NMR and UPLC-MS on the same samples. This method has enabled the characterisation of drug metabolites (xenometabolome) in epidemiology study.⁷⁸ Other recent statistical spectroscopic methods include statistical homogeneous cluster spectroscopy (SHOCSY)⁷⁹ and automatic spectroscopic data categorisation by clustering analysis (ASCLAN).⁸⁰ These methods aim to remove the influence of irrelevant interferences within the dataset to enhance the biomarkers selection process,⁷⁹ and to reliably differentiate potential discriminatory markers from non-discriminatory markers in a biological dataset,⁸⁰ respectively. Figure 2 shows the steps of statistical analysis of INTERMAP NMR metabolic phenotyping data.^{13, 81, 82}

In MWA studies, hundreds to thousands of biomarkers are assayed leading to data that are highly multivariate and co-linear. In order to detect statistically significant relationships between molecular variables and phenotype, we defined the metabolome-wide significance level (MWSL), a threshold required to control the family wise error rate (FWER) through a permutation approach.⁸³ Using the spectra of the INTERMAP Chinese participants (N=836) as the reference population, we investigated the influence of spectral resolution and the number of variables in the NMR spectra, and examined population heterogeneity by repeating the analysis in the INTERMAP US population samples (N=2,164). Results showed

that MWSL of 2×10^{-5} and 4×10^{-6} for a FWER of 0.05 and 0.01 could be used as a benchmark for NMR based MWA studies of human urine. For the subsequent INTERMAP MWA studies, MWSL of 4.0×10^{-6} is used to identify significant metabolic features, as a conservative estimate taking account of the high degree of co-linearity in urinary NMR spectral data.^{81, 82} A similar approach may be used for MS based MWA studies.

Novel biomarker discovery

Identification of unknown discriminatory metabolite is a key bottleneck in MWA study since the human metabolomes is still largely unknown. Elucidating the chemical structure of unknown metabolite is often labour-intensive and requires multiple analyses using a series of analytical experiment. Within the INTERMAP, we have been successful in identification of a number of metabolites including metabolites derived from dietary and drug intake.⁸⁴⁻⁸⁷

Ethyl glucoside

The INTERMAP MWA study has discovered novel metabolite related to the intake of alcohol amongst the Chinese and Japanese population samples. From the ^1H NMR urinary spectra, a doublet at $\delta 4.93$ corresponding to ethyl glucoside was observed and it was derived following the ingestion of rice wine and sake.⁸⁴ This doublet was not observed amongst the western population samples in the INTERMAP Study.

Proline betaine

We used metabolic phenotyping by ^1H NMR spectroscopy to detect an increased excretion of proline betaine, tartaric acid and hippurate after fruit consumption compared with baseline diet in a dietary intervention study.⁸⁵ We then measured concentrations of proline betaine in selected fruit and commercially available fruit juices by ^1H NMR spectroscopy optimized for quantification of this compound. All citrus fruit tested contained proline betaine; concentrations varied from 75 mg/L in orange squash to 1,316 mg/L in orange juice from concentrate. After consumption of 250 ml orange juice we found a singlet peak at $\delta 3.11$, representing the CH_3 moiety of proline betaine. Most proline betaine excretion occurred in the first 14 hours after consumption, peaked at the 2-hr post-intervention urine collection and declined to almost baseline level after 24 hours.

The 24-h dietary recalls of INTERMAP UK participants were then assessed to validate the use of proline betaine excretion as a biomarker of citrus fruit consumption and as a surrogate marker of healthier eating patterns.⁸⁵ Proline betaine excretion differed significantly between individuals with no recorded citrus consumption in their 24-h dietary recall data and individuals with recorded citrus consumption ($P < 0.0001$). Those who reported citrus consumption and had higher levels of proline betaine excretion also showed a healthier nutrient profile with higher intake of vegetable protein, and lower intakes of total fat, trans fatty acids, cholesterol and animal protein, lower urinary sodium-potassium (Na/K) ratio, lower body mass index (BMI) and lower systolic BP compared with non-citrus consumers. These findings provide proof that metabolic phenotyping can discover novel dietary biomarkers that can be used to validate dietary assessment in large-scale epidemiologic data.

Acetaminophen and ibuprofen

Although the metabolism of commonly used analgesics such as acetaminophen and ibuprofen has been widely described^{40, 41, 88, 89}, within the INTERMAP Study, we showed that we could detect urinary metabolite signatures related to commonly used analgesics such as acetaminophen and ibuprofen,⁸⁶ enabling the use of metabolic signatures to verify the self-reported data.⁸⁷ We applied PCA on ¹H NMR spectra of US participants and identified 413 urine samples containing acetaminophen and its metabolites (acetaminophen users) and then applied OPLS-DA analysis on a subset of 70 urine samples from acetaminophen users and 70 urine samples from acetaminophen non-users.⁸⁶ The OPLS-DA loading coefficient plot showed that differentiation between acetaminophen users vs. non-users primarily resulted from the presence of acetaminophen and its metabolites acetaminophen glucuronide, acetaminophen sulfate, and the acetylcysteine conjugate of acetaminophen. Similarly, a PCA model was constructed to identify urine samples of ibuprofen users, and OPLS-DA was performed on a subset of urine samples. The OPLS-DA loading coefficient plot showed that participants who had ingested ibuprofen were differentiated from non-users by the presence of 2-hydroxy, carboxy, and glucuronide conjugates of ibuprofen in the urinary NMR spectra.

We then used these metabolic signatures to verify self-reported analgesic use of INTERMAP participants.⁸⁷ Urinary spectra of UK (Belfast, N=216) and US (Chicago, N=280) participants were inspected for the presence or absence of metabolites in spectral regions containing acetaminophen and ibuprofen.⁸⁷ These spectra were used to construct prediction

models (sensitivity > 98%) based on self-reported analgesic use; overall rates of concordance between questionnaire data and urinary spectra were high for both populations: 83.8% (95% confidence interval, CI: 78.9, 88.7) in Belfast and 81.1% (95% CI: 76.5, 85.7) in Chicago. Overall rates of under-detection of acetaminophen and ibuprofen were low (about 1%) and were comparable for both Belfast and Chicago. We then applied these prediction models to 9,260 urine spectra to evaluate reported analgesic use from self-reported questionnaire. High-level concordance was observed between self-reported analgesic use and ¹H NMR-detected urinary acetaminophen and/or ibuprofen metabolites for all Western population samples, an overall concordance of 70.5% (95% CI: 68.7, 72.2). Our findings demonstrated the efficacy of an objective ¹H NMR-based method for validation of self-reported data on analgesic use, detecting an underreporting rate of approximately 15% in the INTERMAP Study. This MWA approach has demonstrated the potential of metabolic phenotyping in reducing recall bias and other biases in epidemiologic studies for a range of substances, including pharmaceuticals, dietary supplements, and foods.

INTERMAP Metabolome-Wide Association Study

INTERMAP is the first large-scale human population MWA study on diet and BP, using an exploratory analytical approach to investigate metabolic phenotype variation across and within 17 population samples in East (China and Japan) and West (UK and US) based on ¹H NMR spectroscopy.¹³ Using a hierarchical clustering algorithm, we investigated similarity/dissimilarity between populations based on their urinary profiles. East Asian and Western populations had well-differentiated metabolic phenotypes (Figure 3). Among the East Asian samples, Japan was differentiated from China, and within China, North China was differentiated from South despite similar genetic background. Using O-PLS-DA^{70, 73} we reported significant differences of metabolic profiles among these populations; discriminatory metabolites included gut microbial-host co-metabolites (hippurate, PAG and methylamines), amino acids (alanine, lysine, taurine), dietary related metabolites (e.g., ethyl glucoside, trimethylamine-*N*-oxide), compounds related to energy metabolism (acetylcarnitine), and tricarboxylic acid cycle intermediates (succinate and citrate). Four discriminatory metabolites reflecting diet and gut microbial activities, alanine, formate, hippurate and *N*-methylnicotinate (NMNA), were then quantified from the ¹H NMR urinary spectral profiles. We found that alanine was highly correlated with 2-oxoglutarate (metabolic

linkage via glutamate-pyruvate transaminase activity) and with formate (pyruvate/Co-A metabolism), and hippurate was highly correlated with NMNA (renal transporter/secretion mechanisms). In multiple linear regression models, both formate and hippurate were inversely associated with systolic and diastolic BP, and alanine was positively associated with BP.

More detailed analysis was later performed on the Chinese population samples.⁸¹ We found that urinary metabolites significantly different between northern and southern Chinese, reflecting the variations in dietary pattern as well as CVD risk between these two populations; higher in northern than southern Chinese populations included dimethylglycine, alanine, lactate, branched-chain amino acids (isoleucine, leucine, valine), N-acetyls of glycoprotein fragments (including uromodulin), N-acetyl neuraminic acid, pentanoic/heptanoic acid, and methylguanidine; metabolites significantly higher in the south included gut microbial-host co-metabolites (hippurate, 4-cresyl sulfate, PAG, 2-hydroxyisobutyrate), succinate, creatine, scyllo-inositol, proline betaine, and trans-aconitate. Compared to the south, northern Chinese had higher BMI, less favourable diet including lower Ca, Mg and P intakes, higher 24-h urinary Na excretion, higher urinary Na/K ratio excretion, and higher BP (**Table 1**).⁹⁰ The significant north-south differences in BP, BMI and diet⁹⁰⁻⁹² are reflected in geographic variations in both CVD incidence and mortality rates, with higher rates in the north than the south.⁹³⁻⁹⁵ The INTERMAP MWA study indicates the likely importance of environmental influences (e.g., diet), endogenous metabolism, and mammalian-gut microbial co-metabolism, in helping to explain north-south China differences in CVD risk.

The INTERMAP Study confirmed that African Americans (AA) had higher systolic and diastolic BP compared to non-Hispanic white Americans (NHWA)⁸² and this BP difference, in part, was due to less favourable multiple nutrient intake by AA, with lower intakes of fruits, vegetables, and dairy products, and lower intakes of vegetable protein, starch, fibre, K, Ca, Mg, P compared with that of whites. In addition, there was greater obesity prevalence among black compared with white women.^{82 96} ¹H NMR spectra of the INTERMAP US participants showed that urinary metabolites significantly higher in AA than in NHWA included creatinine, 3-hydroxyisovalerate, N-acetyls of glycoprotein fragments, dimethylglycine, lysine, N-acetyl neuraminic acid, leucine, dimethylamine, taurine, and 2-hydroxy-isobutyrate; metabolites significantly higher in NHWA included trimethylamine, NMNA, hippurate, and succinate

(Figure 4).⁸² The mean values of urinary hippurate (2.9 mmol/24-h for AA men, vs. 4.1 mmol/24-h for NHTWA men, $P < 5 \times 10^{-9}$) and NMNA (0.24 mmol/24-h for AA men, vs. 0.42 mmol/24-h for NHTWA men, $P < 3 \times 10^{-10}$) were significantly lower in AA compared to NHTWA. Multiple linear regression was used to examine these AA-NHTWA differences in dietary and urinary metabolites in relation to BP; multiple foods, nutrients and metabolites accounted for part of the higher BP among AA.

Summary

Over recent years, major advances have been made in the metabolic phenotyping of epidemiological samples. The advancement in the analytical techniques and the development of new statistical data analysis tools have enabled the identification of novel metabolic phenotypes associated with diet (including Na intake), xenobiotics and BP. The findings of the INTERMAP MWA study may provide insights into molecular pathways underlying complex biological processes such as adverse BP levels. We envisage that future studies will include the generation of testable hypotheses based on the findings from the INTERMAP MWA study. Moreover, the increasing number of population based cohort studies, which also apply MWA approach, will undoubtedly contribute to our understanding on the key mechanism that are associated with CVD. As noted above, metabolomic data with hundreds to thousands of biomarkers being assayed, are highly multivariate, co-linear and noisy, with potential for false positive findings; it is also always a possibility but unlikely that phenotypes are specific to INTERMAP Study populations and not generalizable; replication studies are needed. Nonetheless, it is reasonable to state at this juncture that INTERMAP findings to date have demonstrated significant independent relationships of several nutrients/foods/eating patterns/metabolites to BP, thereby moving the field forward in exciting and unprecedented ways.

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Figure Legend

Figure 1 INTERMAP metabolome-wide association (MWA) studies.^{12, 13}

Figure 2 Steps involved in the statistical analysis of INTERMAP NMR metabolic phenotyping data.^{13, 81, 82}

Figure 3 Hierarchical cluster analysis (HCA) on ¹H NMR urine spectra, the INTERMAP Study.¹³ The HCA algorithm produces a dendrogram showing the overall similarity/dissimilarity between population samples. Similarity index is normalized to intercluster distance. Each branch of the dendrogram defines a subcluster; population samples within subclusters are more similar to each other than to those in other subclusters. The dendrogram shows clustering based on country, and geographical location or gender.

Figure 4 The median urinary ¹H NMR spectrum of African Americans and non-Hispanic white Americans.⁸² Top: median urinary ¹H NMR spectrum of INTERMAP U.S. AA and NHWA participants, based on the first urine collection (N=1,455). Bottom: Manhattan plot indicating the significant spectral variables. Metabolites higher in AA individuals compared to NHWA are shown in red; in blue for metabolites higher in NHWA individuals compared to AA. Key: 1, Leucine; 2, 3-hydroxyisovalerate; 3, 2-hydroxyisobutyrate; 4, N-acetyls of glycoprotein fragments; 5, N-acetyl neuraminic acid; 6, Succinate; 7, Dimethylamine; 8, Trimethylamine; 9, Dimethylglycine; 10, Lysine; 11, Creatinine; 12, Hippurate; 13, N-methyl nicotinic acid.