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USING METABOLOMICS FOR THE EXPLORATION OF DIET AND HEALTH IN TWINS

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Thesis submitted in fulfilment of the requirements for the degree of Doctor of Philosophy

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Abstract

Metabolomics is an exciting area of research obtaining the chemical and metabolic signals from biosamples. Food consumption is strongly linked to metabolism making the metabolome an ideal phenotype for identifying biomarkers and helping refine and explore diet-disease associations. However, the metabolome is highly complex and influenced by many factors, such as age, disease, genetics and gut microbiota. Discordant monozygotic twins may provide a strong model for confirming association findings as they are matched for age, sex and the baseline genetic sequence.

In this thesis, I explored the potential and applicability of the metabolome in nutritional research in two main areas: for identifying biomarkers of food exposure and further investigating the relationship of food intake with indicators of health. Firstly, I examined metabolomics profiles associated with self-reported food intakes and dietary patterns and confirmed these associations using the co-twin control method. I then identified top metabolite markers of food group intakes, and created and tested metabolite scores using multiple-methods. I searched for markers of gut microbiome diversity, an emerging indicator of health, by examining the association of alpha-diversity with blood metabolomics profiles and the relationship with diet and the metabolic syndrome. In the final chapter, I created a dietary score predictive of visceral fat mass, a strong risk factor for cardio-metabolic disease, and examined the degree to which the relationship between diet and visceral fat mass is mediated by associated metabolites and microbiome taxa. Throughout each chapter I used discordant monozygotic twins to validate top results. Overall, my findings show that metabolomics is a highly versatile tool for advancing nutrition research from biomarker identification through to untangling the impact of dietary exposures on indicators of metabolic health and the gut microbiome.

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Abbreviations

Abbreviation	Meaning
ACSM2A	acyl-CoA synthetase medium-chain family member 2A
ACSM5	acyl-CoA synthetase medium-chain family member 5
AHR	Aryl hydrocarbon receptor
AUC	Area under the receiver operating characteristic curve
BCAA	branched-chain amino acids
BMI	Body mass index
CMPF	3-carboxy-4-methyl-5-propyl-2-furanpropanoate
CPS1	Carbamoyl-phosphate synthase 1
CVD	Cardiovascular disease
CYP1A1	Cytochrome P450 family 1 subfamily A polypeptide 1
CYP1A2	Cytochrome P450 family 1 subfamily A member 2
DHA	Docosahexaenoate
DHEA	Dehydroepiandrosterone
DNA	Deoxyribonucleic acid
DXA	Dual-energy X-ray absorptiometry
DZ	Dizogotic
EFA	Essential fatty acid
EPA	Eicosapentaenoate
EPIC	European Prospective Investigation into Diet and Cancer
F&V	Fruit and vegetable diet pattern
FA	Fatty acid
FADS1	Fatty acid desaturase 1
FBG	Fasting plasma glucose
FD	Fermented dairy
FF	Fried and fast foods
FFQ	Food frequency questionnaire
GAT2	gamma-Aminobutyric acid transporter 2
GWAS	Genome-wide association study
HAO2	Hydroxyacid oxidase 2
HDL	High density lipoprotein cholesterol
KORA	Cooperative health research in the Region of Augsburg
MACS	Medium-chain acyl-CoA synthetase
MDS	Mediterranean diet score
MetS	Metabolic syndrome
MUFA	Monounsaturated fatty acid
MZ	Monozygotic
NAFLD	Non-alcoholic fatty liver disease
NASH	Non-alcoholic steatohepatitis
NAT2	N-acetyltransferase 2
NHANES	National Health and Nutrition Examination Survey
NS	Not significant
NSP	Non-starch polysaccharide
Ost-alpha	Organic solute transporter alpha
Ost-beta	Organic solute transporter beta
OTU	Operational taxonomic unit
PCA	Principal components analysis

PPARα	Peroxisome proliferator-activated receptor alpha
ΡΡΑRγ	Peroxisome proliferator-activated receptor gamma
PUFA	Polyunsaturated fatty acid
ROC	Receiver operating characteristic curve
SD	Standard deviation
SE	Standard error
SLC16A9	Solute carrier family 16 member 9
SLC51A	Solute carrier family 51 alpha subunit
SLC5A11	Solute carrier family 5 member 11
SLC6A13	Solute carrier family 6 member 13
SLC6A20	Solute carrier family 6 member 20
SLE	Systemic lupus erythematosus
SNP	Single nucleotide polymorphism
SSB	Sugar-sweetened beverage
SULT2A1	Sulfotransferase family 2A member 1
TG	Triglycerides
ТМАО	Trimethylamine N-oxide
VF	Visceral fat
VFM	Visceral fat mass
WC	Waist circumference
WDR85	diphthamide biosynthesis 7
WG	Whole grain

Chapter 1 Introduction

In this chapter I discuss twin studies that have examined the heritability of reported food and nutrient intakes and dietary patterns. I then introduce the usage of monozygotic (MZ) twins for recent nutriomics studies including specifically the metabolome and gut microbiome. Finally, I briefly introduce the usage of twins combined with nutriomic methods for the exploration of the effect of diet on metabolic disease.

Part of this work has been published as a literature review in *Nutrition Research Reviews* (Pallister et al., 2014).

Early nutrition studies on twins and multi-foetal pregnancies focused on the side effects of the pregnancy, achieving adequate maternal nutrition during gestation, and on the implications for the health of the growing foetuses (Brown and Carlson, 2000). The womb, however, is a solitary internal environment, where nutritional parameters can be measured with more ease. Once twins enter the world, where myriad of external environmental influences on different phenotypes can be assessed, twins provide us with the unique ability to accurately determine heritability of certain complex traits, such as dietary phenotypes. In this context, dietary phenotypes broadly refer to the observable and measurable foods, and their constituents, consumed by individuals or groups. Indeed, twin studies have proven that genetic makeup plays a significant role in dietary intakes (Rankinen and Bouchard, 2006). Moreover, twins have the advantage of limiting inter-individual variability as they are raised in a similar environment and have matched age, genes and sex (for monozygotic [MZ] pairs). They therefore represent a natural matched case-control experiment.

Omics technologies allow for a systems approach to nutrition research, that encompasses the primary level of DNA sequence (genomics), gene expression (transcriptomics) and epigenome, to the intermediate phenotypes (proteome, metabolome, and microbiome), finally to clinical endpoints (Kussmann et al., 2006). Food constituents may influence each of these areas, but their effect is difficult to ascertain due to imprecise dietary assessment methods. Omics techniques may help ameliorate this issue through food intake biomarker discovery within the intermediate phenotypes, primarily the metabolome (Zivkovic and German, 2009). Once biomarkers of intake

have been validated, in conjunction with currently utilized intake and known biomarker measures, reliable associations between genomic and transcriptomic data can be made. This can allow for identification of groups at risk for poor dietary patterns, gene-diet interactions, and diet-disease associations, which combined could conjure an accurate depiction of a "healthy" phenotype (Kussmann et al., 2006), although this is limited by currently used methods of dietary assessment.

Presently, epidemiological studies rely mainly on subjective reporting to obtain dietary information. These include prospective methods such as weighted and estimated food records, and retrospective methods, including 24-hour recalls and food frequency questionnaires (FFQs). Each of these methods carry their own strengths and weaknesses, which have been reviewed by Shim et al. (2014). Weighted records and estimated food records carry a higher participant burden, and use more time and costs, though they provide more accurate data as they do not rely on participant memory (i.e. recall bias). 24 hour recalls commonly require an interviewer (though computer-based methods are becoming more common-place), they are limited by possible recall bias and interviewer bias and are expensive and time-consuming. Both food records and 24 hour recalls require multiple days of evaluation to assess usual food intake. FFQs are the most commonly used dietary assessment method for population-based studies as they have a lower participant burden and resources and indicate habitual food consumption, though are limited to a set list of foods and have low accuracy due to recall bias. With these limitations in mind, identifying novel dietary biomarkers using multi-omics will be challenging and require confirmation and evaluation by dietary intervention studies.

In this introduction, I first discuss the usage of twins for generating heritability estimates of food intake phenotypes, including energy and macronutrient intakes, dietary patterns and specific food group intakes. I then highlight the value of discordant MZ pairs, for furthering nutrition research through discovery and validation of biomarkers of intake and health status in conjunction with cutting-edge omics technologies, with metabolomics being the focal point.

1.1 Heritability of food intake phenotypes: The usage of twins for heritability estimates of intakes

The classic twin design allows for a natural experiment that exploits the difference in genetic relatedness between MZ and dizygotic (DZ) twins to estimate the degree to which phenotypic variability, such as food or nutrient intake, is explained by genetic and environmental factors. Heritability is defined as the degree of total phenotypic variance due to genetic variation and is relevant only to groups or populations and not at the individual level (Visscher et al., 2008).

1.1.1 Heritability of energy and macronutrient intakes

Accurate assessments of the genetic influence on energy intakes is crucial as these provide a baseline assessment for which genes influence total food intakes (de Castro, 1993a) and therefore justify genotype-specific dietary intervention strategies. Furthermore, the degree to which macronutrient intakes are genetically determined may have significant implications for the role of elevated intakes of energy dense foods (i.e. those containing high concentrations of fat and refined carbohydrates) to the obesity epidemic (Guyenet and Schwartz, 2012). Therefore, multiple twin studies have evaluated these components through various dietary assessment methods (**Figure 1-1a: Appendix A Table 1**)

1a; Appendix A Table 1).Studies of different size and quality have estimated heritability of different energy and

macronutrient intakes to lie between 8% and 70% (Liu et al., 2013, Pimpin et al., 2013, de Castro, 1993a, Heller et al., 1988, Wade et al., 1981, Hasselbalch et al., 2008, Aden et al., 1979, Hur et al., 1998). Beginning from a young age, infant twins have showed that the genetic component provides a small, albeit significant, effect on energy and macronutrient intakes (8% to 12%) (Pimpin et al., 2013⁾. As children age and they become less dependent on their parents, the genetic effect on energy and macronutrient intakes appears to intensify, as suggested by findings in 11-13 year-old US twins (31% to 48%) (Liu et al., 2013). This pattern is similar to other health-related phenotypes, including BMI (Dubois et al., 2012), which has been recently confirmed through a more complex longitudinal genome-wide complex trait analysis for determining DNA-based heritability (Llewellyn et al., 2014). Heritability of dietary energy and macronutrients in adults has been found to vary widely: for energy (32% to 65%), for fats (35% to 53%), carbohydrates (25% to 67%), and proteins (28% to

70%) (Hasselbalch et al., 2008, Hur et al., 1998, de Castro, 1993a, Wade et al., 1981). An extensive study of adult Danish twins showed inheritance of energy and energy-adjusted macronutrient intakes to range from 28% to 55% (Hasselbalch et al., 2008). Although, genetic effects on macronutrient intakes were more substantial in men (49% to 55%) than women (28% to 36%), they were not different between macronutrients. In support of this, two other twin studies suggested the same genetic mechanisms governing energy intakes (Guyenet and Schwartz, 2012) influence macronutrient intakes (de Castro, 1993b, Faith et al., 1999).

The lack of evidence to date for independent sets of genes influencing macronutrient intakes from those genes governing energy intakes has focussed attention on whether dietary energy density (kilocalories per gram of food) is genetically determined, with one study suggesting substantial genetic influence (de Castro, 2006), and another not (Hasselbalch et al., 2008). Despite this, in the latter Danish study, factors related to dietary energy density, including fibre, glycaemic index and the glycaemic load, were found to be significantly heritable both in women (49%, 36% and 33%, respectively) and men (41%, 30%, and 25%, respectively). The dependence of these phenotypes on the constituents of the whole diet suggests genetic effects on these phenotypes may be reflective of variable dietary patterns.

1.1.2 Heritability of dietary patterns

Dietary patterns are identified by two primary methods: empirically-derived or a priori (Hu, 2002). Empirically-derived dietary patterns employ statistical methods to identify natural groupings of intakes of food items (e.g., principal component analysis or cluster analysis (Hu, 2002)). A priori dietary patterns are generated based on adherence to diet parameters previously associated with health outcomes or physiological states, such as the Mediterranean diet score. A key advantage to using dietary patterns is the translatable results that encompass the degree to which genetics influence the global diet.

Two twin studies on children using empirically-derived dietary patterns generated highly inconsistent heritability estimates despite reasonable sample size (**Appendix A Table 2**) (Breen et al., 2006, Faith et al., 2008). In a study of US 7 year olds (*n*=792) heritability estimates of food pattern intakes obtained by 24-hour recalls were stronger in boys than girls, ranging from 12% to 79% and 20% to 56%, respectively (Faith et al., 2008). In another study of 4 to 5 (SD: 0.3) year-old

UK twin-pairs (*n*=214) heritability of food preference was measured through the use of a modified 95-item "liking" food frequency questionnaire (FFQ) administered to parents (Breen et al., 2006). Heritability ranged from 20% (desserts) to 78% (meat and fish) on food liking-disliking groups. It is difficult to make valid comparisons between these two studies as the dietary assessment methods used measure different outcomes (i.e. intakes versus liking), although it has been found that exposures to foods generally encourages liking and intakes in children (Wardle et al., 2001). This suggests environmental influences may override genetic predispositions to food pattern intakes in children, which has implications for combating food neophobia, a highly heritable and perhaps evolutionarily important trait (Cooke et al., 2007).

Contrary to studies in children, studies in adults have generated relatively stable heritability estimates for food pattern intakes across patterns and genders. "Healthy" dietary patterns, characterised by high intake of vegetables, fruits and whole grains and low intake of fatty foods of animal origin and simple carbohydrates, have heritability estimates ranging from 33% to 54%, while "unhealthy" patterns ranged from 33% to 50% (Appendix A Table 2) (van den Berg et al., 2013, Keskitalo et al., 2008, Teucher et al., 2007, Gunderson et al., 2006, van den Bree et al., 1999). Furthermore, a study which utilized a twins of mistaken zygosity approach (a useful method to control for potential equal environmental assumption bias (Scarr, 1968)) in US female twins (n=700), found additive genetic effects to account for 50% of the variability in healthy pattern intakes, while an unhealthy diet pattern was not significantly heritable (Gunderson et al., 2006). A healthy diet is more phenotypically similar to diets humans have consumed through millennia and evolved with, it is only very recently that high energy dense and processed foods have been readily available, as such it makes sense the variation in this diet may be more genetically influenced in some groups (Breslin, 2013). One study conducted on an older age group (≥ 50 years) determined additive genetic effects to account for 33% of the variability of consumption frequency for both healthy and unhealthy patterns (van den Bree et al., 1999). A pattern of increased dietary variety in modern societies may contribute to the obesity epidemic through ready availability of highly processed, calorific foods compared to healthier, natural choices such as fruits and vegetables (McCrory et al., 1999). In middle age to elderly twins from the "Virginia 30,000" twin study (N = 5,543) dietary variety seeking was up to 30% heritable, supporting the notion that our environment is an important driver of dietary variety (Scheibehenne et al., 2014).



Figure 1-1. Estimated heritability of nutrients (a), food (b) and beverage (c) intakes from twin studies

Intake heritabilities presented are significant findings from previous twin studies. Heritability histograms are color coded according to study. From clockwise, histograms are grouped according to age; the first line below the histogram denotes this: light grey, children; dark grey, adults. Within each age group, histograms were grouped according to accuracy of the dietary assessment method used, from most accurate (e.g. 2 day buffet style meal intervention) to least accurate (e.g. 67-item FFQ). The second line below the histograms indicates gender: pink, female; blue, male; yellow, combined. *Modified from Pallister et al. (Pallister et al., 2014)*

1.1.3 Individual foods and food group heritabilities

A number of recent studies have estimated the genetic effect of food and beverage intakes in humans (**Figures 1-1b** and **1-1c**; **Appendix A Tables 3** and **4**) generating wide-ranging heritability estimates across food groups, particularly for children (Pimpin et al., 2013, Fildes et al., 2014). Findings from the UK Gemini cohort of infants and children aged 21 months to 3 years have shown age effects on heritability of food group intakes are evident even in the early years (Fildes et al., 2014, Pimpin et al., 2013). Intakes for 21 month-old children were minimally heritable for food groups ranging from 9% (potatoes) to 17% (dairy). Whereas preference for food groups were highly heritable in 3 year-old children, ranging from 27% (dairy) to 54% (vegetables). It should be noted that heritability estimates in the older children were calculated from age- and sex-adjusted residuals, an approach which may have prevented inflation of shared environmental effects (McGue and Bouchard, 1984). The study on infants did not use this approach, perhaps resulting in an underestimation of the genetic effects on food group intakes.

Recent studies in adults have also confirmed there is a genetic component to many food type intakes. Danish twins showed a large degree of variation in calculated heritabilities overall ranging from 17% (fish) to 68% (potato) in men, and 20% (whole grain cereal) to 61% (fish) in women (Hasselbalch et al., 2008). Similar results were seen in UK female twins where heritabilities ranged from 8% (refined grains) to 46% (garlic) (Teucher et al., 2007). Furthermore, the grouping of fruit and vegetable sources generated the highest heritability estimate in this cohort (49%). These authors proposed taste perception may be a key driver of the predicted genetic influence as foods characterised by strong tastes are the cornerstone of these groups. This observation extends to consistently substantial heritabilities for coffee as well as alcohol intakes, ranging from 29% to 73% and 28% to 82%, respectively (Hasselbalch et al., 2008, Teucher et al., 2007, Hur et al., 1998, de Castro, 1993b). Although, the pharmacological properties of both must be considered as alcohol dependence, as well as caffeine consumption-related traits, are both highly heritable (30% to 70%, and 36% to 58%, respectively) (Agrawal and Lynskey, 2008, Yang et al., 2010a) and may drive intakes up in a way independent of taste preference or a biological requirement for these items.

Taste is the strongest determinant of food choices (Feeney et al., 2011), and a recent twin study showed multiple chemosensory facets of numerous food compounds to be highly heritable

and associated with particular gene variants (Knaapila et al., 2012). Future heritability studies of food intakes may shift the focus to food groupings characterized by specific tastes.

1.1.4 Nutritional phenotype heritability conclusion

In summary, twin studies have shown food and macronutrient intakes to be genetically determined. However, heritability estimates are often inconsistent between studies for multiple reasons. First, heritability is only specific to the population studied, which can vary greatly, due to the vast country specific food environments. Indeed, if a food is not readily available in a population there is no capacity for intakes to be genetically determined. Second, important evidence is available for familial and community influence on food intake (such as parenting styles, food exposures, socioeconomic status) which have an important place for battling current issues, such as childhood obesity (Berge, 2009) and food neophobia (Anzman-Frasca et al., 2012), and diabetes (Pollard et al., 2014). Finally, methods for determining heritability are not standardised, for instance the inclusion/exclusion of variables from the model (e.g. age, country), the segregation/aggregation of sexes, and the different methods for obtaining and analysing dietary data tend to be unique for each study. The same genetic mechanisms governing energy intakes are suspected to influence macronutrient intakes. However, recent studies have assessed the impact of gene variants encoding proteins related to the complex energy intake regulatory systems under central nervous system control (Guyenet and Schwartz, 2012) on preference for macronutrient types. These studies are few on twins (Hasselbalch et al., 2010, Bouchard-Mercier et al., 2012). Heritability estimates are more limited for macronutrient subtype intakes (e.g. the fatty acid profile (Hur et al., 1998, Heller et al., 1988)), different age groups or changing physiological states (e.g. pregnancy), and socially driven diets (e.g. veganism). Macronutrient profiles do not account entirely for food pattern preferences that may be more determined by genetic makeup. Variation in diet patterns of adults are to a relatively high degree attributable to genetics (van den Berg et al., 2013, Keskitalo et al., 2008, Teucher et al., 2007, Gunderson et al., 2006, van den Bree et al., 1999). Furthermore, specific food type intakes that characterise a healthy diet pattern tend to be highly heritable in adults and in turn consist of foods that elicit distinctive taste responses (e.g. garlic, fruit and vegetables) (Teucher et al., 2007).

The substantial heritability of multiple dietary phenotypes supports the concept of personalised dietary recommendations, the generation of which has so far primarily relied on the exploration of an individual's genetic makeup. However, on a molecular level, the search for genotypes predisposing individuals to particular dietary intakes is very much in its infancy. Nonetheless, the evidence generated by twin heritability studies can lead scientists to dietary phenotypes that are consistently highly heritable and demand further exploration through GWAS and candidate gene studies, following which twins can aid the determination of the heritable contribution of specific singly nucleotide polymorphisms (SNPs) to the dietary phenotype (as used previously for human height (Yang et al., 2010b)).

A key issue which has long plagued nutrigenetic research arises from the inaccuracies of self-reported diet intake and food preference data and as a result, the hunt for dietary biomarkers, objective measures of dietary intakes and nutritional status, has been hastened. These may allow for more accurate heritability calculations for food and nutrient intakes as has recently been shown for salt intakes using urinary sodium excretion, an established, valid biomarker of salt intake (Kho et al., 2013), and could potentially be used in genetic association studies. However, the usage of biomarkers is not confined to genetic studies. It is becoming increasingly clear that for the establishment and refinement of personalised nutrition, in combination with genetic information, a whole organism, multi-systems approach must be undertaken, where the usage of biomarkers of food intakes and nutritional status will be essential. Nutrition research is now embarking on a nutriomics era, where emerging high-throughput omics technologies will substantially aid biomarker discovery and "healthy phenotype" definition. As will be explored next, twin studies are progressively taking steps to make a contribution in this way.

1.2 The importance of twins for emerging nutriomic research

This next section will explore emerging dietary studies on twins used in collaboration with highthroughput omics to capture metabolism (metabolomics) and the gut microbiota (microbiome). Integrating these technologies to determine a "healthy" phenotype is the long-term goal of personalised nutrition (van Ommen and Stierum, 2002). A reoccurring issue with this type of analysis is the high degree of inter-individual variation, making associations often inconsistent (Kussmann et al., 2006). As such, the usage of identical twins discordant for dietary factors or

objective measures of nutrition status, matched entirely for age, sex, genetics and partially for early environmental influences, will provide an enhanced method of assessing diet and biological relationships.

Obesity, the greatest and fastest growing health concern in the world at this time, is a highly genetic disorder, influenced by myriad complex factors (Despres et al., 1992). However, rare obese discordant identical twin pairs, provide a unique opportunity to disentangle lifestyle and environmental factors on the human system derangements induced by energy imbalance independently of genetics.

1.2.1 Nutrimetabolomics

Nutritional metabolomics involves the extensive chemical profiling of various tissues completed in a global manner through targeted and non-targeted methods used as a complement to diet- and health-related complex systems approaches. The metabolome gives unique information into the metabolic status of an individual by providing a snapshot of the metabolic processes undertaken in a bodily system, specific organ, tissue or cell, which cannot be identified through measuring gene expression or the proteome. Targeted methods measure a panel or group of chemically defined metabolites. Whereas non-targeted methods aim to detect all measureable chemicals in a sample and therefore have more coverage. Moreover, non-targeted metabolomics has the potential to detect previously unknown metabolites in a sample, increasing the possibility for identifying novel findings. Although, it may be difficult to draw conclusions from such findings. Targeted methods used for non-targeted methods. Moreover, targeted methods used for non-targeted methods. Moreover, targeted methods typically measure metabolites which are biologically well-defined (Patti et al., 2012).

The two primary technologies used to analyse sample metabolomes are mass spectrometry (MS) and nuclear magnetic resonance (NMR) spectroscopy. NMR spectroscopy is a quantitative technique that determines information on the solution-state molecular structures in a sample through atom-based nuclear interactions (Marion, 2013). MS-based methods typically separate metabolites first in a sample using chromatography (examples include gas or liquid-based chromatography, and capillary electrophoresis), then metabolites are identified with a mass spectrometer (Feng et al., 2008). Although, some MS methods do not require the initial sample

separation step and metabolites can be quantified directly. Gas or liquid chromatography separate metabolites by time, providing detailed chemical information, though this may require complex chemical preparation that can destroy metabolites (Sellick et al., 2010). Moreover, often techniques (such as GC–MS and LC-MS) are combined as different techniques detect different types of metabolites.

Veenstra (2012) previously summarized the advantages and disadvantages of both methods. The key advantage of MS metabolomics is the sensitivity or the ability to measure very small analytes (within the femtomolar to attomolar range) with accuracy. Although the main disadvantage of MS is less precise quantification due to the MS signal intensity being influenced by the method of sample preparation (Veenstra, 2012), with complex sample preparation also being a disadvantage. Conversely, NMR spectroscopy is highly precise due to the peak area of a compound being proportional to the sample concentration of particular identified nuclei (such as ¹H, ¹³C) (Veenstra, 2012). Moreover, NMR techniques require minimal sample preparation, are non-destructive, and provide rapid results with high reproducibility at a low cost (Markley et al., 2016). The primary disadvantage of NMR-based methods is their lack of sensitivity (Veenstra, 2012), it is also non-discriminatory (Markley et al., 2016).

Nutrition and the metabolome are intimately linked in that nutrients and non-nutrient food constituents supply metabolites, however these contributions make the dietary influence on the metabolome all the more difficult to ascertain (Gibney et al., 2005). Despite this, dietary pattern and food intakes generated through self-reported intake data have identified novel biomarkers within the metabolome (O'Sullivan et al., 2011, Altmaier et al., 2011, Guertin et al., 2014b, Zheng et al., 2014), validating this approach for use in epidemiological studies. However, the genetic influence on metabolite levels is wide-ranging (Suhre et al., 2011), therefore novel usage of the twin model provides an ideal method for determining the dietary impact on metabolites, through segregating the non-genetic component. One nutri-metabolomic study conducted on twins further confirmed self-reported intake associations with metabolites (Menni et al., 2013c). Female UK twins completed 131-item FFQs that generated dietary constituents and intake patterns found to associate with 42 metabolite levels in the larger twin population. Monozygotic twins discordant for dietary intakes were then identified and utilized to replicate results, confirming the utility of this method (Zivkovic and German, 2009).

Lipidomics, a branch of metabolomics, has been used to characterize environmental and lifestyle-induced changes to the global serum lipid profile in 14 healthy, young MZ obesitydiscordant twin pairs (Pietilainen et al., 2007). Levels of lysophosphatidylcholines, lipids associated with inflammation (Yang et al., 2005) and atherogenesis (Glass and Witztum, 2001) were found to be elevated in obese co-twins, with concommitant decreases in antioxidant (Wallner and Schmitz, 2011) ether phospholipids. Moreover, despite the young ages of the obese co-twins (24 to 27 years), these lipid profile changes were associated with insulin resistance.

The metabolome is highly complex and inter-individual variation is high for a multitude of reasons (Zulyniak and Mutch, 2011), including age (Menni et al., 2013b), sex (Krumsiek et al., 2015), genetics (Shin et al., 2014), ethnicity (Wikoff et al., 2013) and the gut microbiome (Wikoff et al., 2009), thus factoring solely nutritional intake into nutrimetabolomic studies is problematic and the relative importance of blood and urine levels is unclear. For example, previous analysis on the non-targeted metabolomics TwinsUK dataset showed 22 metabolites to account for 59% of the variance in age (Menni et al., 2013b). In another study conducted on the TwinsUK dataset the heritability of blood metabolite levels ranged from 10-81%, while only 30% of the blood metabolites were not significantly influenced by genetics (Shin et al., 2014). Finally, sex has shown to influence one-third of metabolites, while a network analysis showed certain metabolic pathways to be sexspecific (Krumsiek et al., 2015).

An area of increasing interest for its potential influence on metabolic processes is the gut microbiome. In rodents, the gut microbiota were found to account for 10% of the variability in the plasma metabolome (Wikoff et al., 2009). Although estimates for humans are unknown at this time, regional variation in human metabolomic profiles has been attributed in part to gut microbes (Nicholson et al., 2012). In turn, the gut microbiota carry out processes in the digestion and handling of nutrients, some of which are essential for optimal nutrition, prior to entering the host metabolome (Nicholson et al., 2012). Emerging methods for direct measurement of the intestinal metabolome will help to unravel the microbiome effects on host metabolome (Ursell et al., 2014).

1.2.2 Microbiome

The human gut microbiome is estimated to contain hundreds of bacterial species, with upwards of 536,000 bacterial genes among them (Qin et al., 2010). For years, dietary constituents, such as

pre- and pro-biotics, have been consumed for their proposed health benefits postulated to be through modulation of the gut microbial population (Collins and Gibson, 1999). Population-based studies suggest dietary patterns appear to strongly drive gut microbiome composition. Differences have been observed between Americans and Malawians and Amerindians (Yatsunenko et al., 2012), and consumers of meat- versus plant-based diets (Matijasic et al., 2013, Wu et al., 2011a), findings that are being confirmed by dietary intervention studies (David et al., 2014, O'Keefe et al., 2015). Although the potential for the gut microbiota to drive eating behaviours has also been proposed through modifying cravings and inducing dysphoria (Alcock et al., 2014), for instance individuals showing a desire for chocolate have a distinctive urinary microbial metabolite profile than subjects who were chocolate indifferent, although they consumed identical diets (Rezzi et al., 2007), studies in this area are limited. Twins discordant for obesity have shown significantly different gut microbial species composition (Tims et al., 2013). Lower BMI was associated with increased primary fibre degraders, while higher BMI subjects displayed an abundant network of butyrate producers, as such it was suggested that a shift in fermentation patterns near the end of the colon may be influencing energy homeostasis, although this study did not investigate the dietary impact on the microbial species population.

An analysis on Finnish MZ twin pairs concordant (*n*=9) and discordant (*n*=11) for BMI revealed dietary energy intakes and constituents to be more influential on microbial populations than BMI group (Simoes et al., 2013). Specifically, stool bacterial counts were significantly influenced by intakes of energy, monounsaturated and *n*-3 and *n*-6 polyunsaturated fatty acids, and soluble fibre intakes. Furthermore, profiles of *Bacteroides* spp. were very similar in co-twins consuming the same amount of energy or saturated fats than twins discordant for intakes. These findings suggest that the obesity status *per se* is less influential on microbial species population and rather, it is the content of the diet (i.e. high energy and fat) mediating important changes, highlighting the essential need to incorporate dietary variables into obesity research. The role of host genetics on the microbiome composition is still unclear, though our group has recently shown that gut microbiome phenotypes ranged from 0% to 39% heritable (Goodrich et al., 2014a), further studies (which are ongoing) are needed to provide clarity.

Recent findings suggest that the microbial species signature has the potential to promote and aggravate an obesity phenotype and may interact synergistically through diet-by-microbiota

interactions. In a unique study, transplantation of fecal samples from US female lean and obese cotwins (3 DZ; 1 MZ) into germ-free mice revealed the obese compared to lean co-twin's microbiota significantly increased adiposity and metabolic derangements in mice (Ridaura et al., 2013). When co-housed with lean mice, obese mice adiposity and metabolic effects were reduced, suspected to be a result of Bacteroidetes translocation from lean to obese mice. When fed a diet based on the lower tertile of saturated fats and upper tertile of fruits and vegetables of the US National Health and Nutrition Examination Survey (NHANES), obese mice adiposity was sustained, but relieved when co-housed with lean. Moreover, when compared to co-housing with normal chow, the healthier diet aided the success of lean to obese mice bacterial invasion. When supplied with an unhealthy diet containing the upper tertile of saturated fats with the lower tertile of fruits and vegetables from the NHANES survey, both lean and obese mice presented with significantly increased adiposity, mitigating transmissible effects. These results show the potentially causal role of the microbiome composition and potential for its manipulation.

1.3 Applying omics to untangle the effects of diet on metabolic disease in twins

Obesity elevates risk of cardiometabolic diseases such as the metabolic syndrome and CVD. The metabolic syndrome is a cluster of metabolic abnormalities characterised by at least three of the five symptoms: high blood pressure, an unfavourable lipid profile specifically elevated triglycerides and reduced high density lipoprotein cholesterol, elevated waist circumference, and elevated fasting blood glucose. One of the primary risk factors for cardiometabolic disease is elevated visceral fat mass (St-Pierre et al., 2002, Naukkarinen et al., 2014), which is metabolically active fat that covers the internal abdominal organs.

In the previous sections I have discussed the independent applications of metabolomics and the microbiome to examining diet, primarily through studies on body weight discordant MZ twins. However the link between diet and metabolic outcomes such as the metabolic syndrome and related factors could potentially be further elucidated through collaborative applications of omics technologies. In one recent twin study (Bogl et al., 2016), using metabolomics in combination with waist circumference (an indicator of visceral fat mass) a strong genetic overlap between waist circumference and a number of metabolites, most notably positive correlations between phenylalanine (rg = 0.40), glycoprotein (rg = 0.37), serum triglycerides (rg = 0.36) and BCAAs (rg = 0.30-0.40), and negative correlations between HDL particle diameter (rg = -0.33) and HDL cholesterol (rg = -0.30), supporting the usage of the MZ discordant twin model.

A single study employed multi-omic methods to examine the time-dependent impact of a Big Mac challenge (Bondia-Pons et al., 2014) on 394 serum metabolites and faecal microbiota in 16 MZ twin pairs discordant for BMI and 9 concordant non-obese MZ pairs (Bondia-Pons et al., 2014). Overall, intra-pair changes in metabolite levels were minimal following the Big Mac challenge. Intrapair differences at baseline for two branched chain amino acid (BCAAs) were found to converge after 120 minutes following the Big Mac challenge, with lower BCAAs in the high weight co-twin compared to the low weight co-twin at baseline. Secondary bile acids glycocholic acid and glycolithocholic acid were significantly different at 120 min following the Big Mac challenge, with higher levels in lean co-twins compared to higher weight co-twins. An examination into Bacteroides spp. diversity did not find significant differences between weight-discordant twins, though overall Bacteroides spp. diversity was positively associated with postprandial changes in sugar- and microbiota-derived organic acids. Links to other clinical indicators of metabolic health were also assessed, in particular postprandial changes in glycine-conjugated ursodeoxycholic acid were negatively associated with liver fat content as well as MI, a marker of insulin sensitivity. This study showed overall that within-pair similarity was the primary determinant of the metabolic response to a dietary challenge, highlighting the important role of genetics and early life factors in metabolism.

1.4 Conclusion

The interconnectedness between each omic presented here is evident. A systems biological approach that incorporates multiple omics methods for phenotype definition for use in nutriomic studies is much needed (Norheim et al., 2012). The principal central to this approach is that the whole organism will provide a more accurate view than the sum of its parts, or rather, the entire system has definitive characteristics which will not be replicated by simply adding the effects (MacLellan et al., 2012). However, a primary setback for amalgamating these methods is the inability to define what is "normal" due to the significant inter-individual variability underlying

metabolic processes (Kussmann et al., 2006). Furthermore, it has been suggested that many omics vary relatively little over time within individuals but significantly between individuals compounded by factors such as age, BMI and gender (Eady et al., 2005). To aid in discerning these effects, longitudinal phenotypic information from twin registries will be a critical resource for molecular dietary studies of the future (van Dongen et al., 2012). The studies presented here fall short of this view of an integrated technology-driven approach and focus rather on omics in isolation as this is where the current research stands. To move forward twin cohorts need to make a collaborative effort, collecting extensive dietary data in conjunction with multiple omics, undertaking complex statistical analyses, while contributing findings to international proposed databases such as the Nutritional Phenotype database (van Ommen et al., 2010).

Identifying individuals susceptible to poor dietary habits and defining an (un)healthy phenotype are the overarching aims of nutrigenetic and nutrigenomic research. Twins have provided valuable evidence that many dietary intakes are influenced by genetics, validating further nutrigenetic research and future dietary counselling which targets this domain. Early studies of twins for nutriomic studies have primarily been for unhealthy phenotype definition and single omic, while being less inclusive of diet. However, future studies need to begin to incorporate dietary factors to fill a void in this research area. Although accurate dietary assessment methods are problematic (Tucker et al., 2013). As twins have shown, metabolites are significantly correlated with dietary constituents, and could in the future be used as surrogates, as well as other techniques (e.g. microbiome) (Lloyd et al., 2013, Vernocchi et al., 2012). Twin studies will remain an important and integral part of nutritional research now and in the future.

2.1 Hypothesis

Metabolites have the potential to act as useful surrogates of dietary intakes. Identification of metabolites in tissues that consistently associate with reported intakes through validation by the cotwin control method can be used to better establish and explore diet-disease relationships.

2.2 Aims

- To identify and characterise metabolite associates of self-reported intakes of foods and dietary patterns.
- To create, validate and test the utility of food group metabolite scores as novel biomarkers of food group intakes.
- iii) To identify metabolite marker(s) of gut microbiome diversity that are modulated by food intake, identify the gut microbiome profile of those markers and their relationship to metabolic disease.
- iv) To create a visceral fat mass dietary risk score and characterise the score using a multiomic approach.

Chapter 3 Materials and methods

In this chapter I provide an overview of the study population used, I briefly describe the food frequency questionnaire used to evaluate food intakes, the metabolomics and genotyping datasets and clinical measures used throughout the thesis. Moreover I introduce the statistical method for calculating heritability.

3.1 Subjects and phenotypes

3.1.1 Subjects

Subjects included in all analyses were twins enrolled on the TwinsUK registry, a sample of extensively phenotyped, mainly female, adult MZ and DZ twins from the UK (Moayyeri et al., 2013, Spector and Williams, 2006). Healthy Caucasian twins were recruited nation-wide primarily through media campaigns. Twin zygosity was determined by a validated questionnaire or through multiplex DNA fingerprinting (PE Applied Biosystems, Foster City, CA). Ethical approval has been obtained from St. Thomas' Hospital Research Ethics committee and all subjects have undergone informed consent. Specifically, my thesis included female and male twins, aged 18 to 80 years, who had completed at least one FFQ between 1994 and 2001 (**Appendix B Document 1**), in 2007, and in 2014 and 2015 (**Appendix B Document 2**).

3.1.2 Food frequency questionnaire

Twins completed a 131-item FFQ that was developed and validated against pre-established nutrient biomarkers for the European Prospective Investigation into Diet and Cancer (EPIC) Norfolk (**Appendix B Documents 1 and 2**) (Bingham et al., 2001, Bingham et al., 1997). Processing of the FFQ, including subject exclusion and determination of nutrient intakes was completed by nutritionists from the University of East Anglia. Intake frequency of an average serving of listed foods was determined from a 9-point scale ranging from "Never or less than once/month" to "6+ per day". The questionnaire was intended to capture average intakes in the past year. Nutrient intakes were determined via consultation with *McCance and Widdowson's The Composition of Foods* 6th

edition (McCance et al., 1991), this resource includes nutritional information (including fat, sodium, fibre, and carbohydrates, vitamins and minerals) for over 1,200 commonly consumed foods derived from the UK Nutrient Databank.

3.1.2.1 Exclusion criteria

Submitted FFQs were excluded if greater than 10 food items were left unanswered, or if the total energy intake estimate derived from FFQ as a ratio of the subject's estimated basal metabolic rate (determined by the Harris-Benedict equation (Frankenfield et al., 1998)) was more than two standard deviations outside the mean of this ratio (< 0.52 or > 2.58). Overall, this included 828 twins (9.4%) and 284 twins (6.5%) who completed FFQs between 1994 and 2007 (*n*=8785) and between 2014 and 2015 (*n*=4400), respectively.

3.1.2.2 Residual energy adjustment

Differences in reported intakes as a product of physiological requirements (due to differences in body size, physical activity, and metabolic efficiency) may mask true diet-outcome associations. To account for this, food and nutrient intakes were energy adjusted using the residual method prior to analysis (Willett and Stampfer, 1986), unless otherwise specified. This involved regressing each food item or nutrient on the estimated total kilocalorie intake. The residual was then added to a constant, in this case, predicted food or nutrient intake for the mean estimated total kilocalorie intake.

3.1.3 Anthropometry

Anthropometric measures including body height (metres) and weight (kilograms) were taken at clinical visits by a trained research assistant. Body mass index (BMI) was calculated as body weight in kilograms divided by the square of height in metres.

3.1.4 Samples

3.1.4.1 Blood samples

Fasted blood samples were collected by a trained research nurse at the twin's annual visit at St. Thomas' Hospital. Twins were instructed to fast for 8 hours prior to their clinical visit. Visits occurred in the morning (9:00 or 10:00) or afternoon (13:00 or 14:00). Time of collection was not recorded

therefore diurnal variation may have influenced the results. Following collection samples were stored at -80°C until further processing.

3.1.4.2 Faecal samples

Faecal samples were collected by the twins at home. Following collection, samples were stored in the refrigerator for 2 days or less prior to their annual clinical visit at St. Thomas' Hospital. Once the samples arrived with the clinical team they were stored at -80° C until further processing.

3.1.5 Clinical measures

3.1.5.1 Blood pressure

Systolic and diastolic blood pressure (SBP and DBP, respectively) were carried out by an experienced nurse using the Marshall mb02 or the Omron Mx3 Digital Blood Pressure Monitors. An average of three readings (separated by one minute) was used for analysis.

3.1.5.2 Blood lipid profiling and glucose

Three devices (Cobas Fara; Roche Diagnostics, Lewes, UK; Kodak Ektachem dry chemistry analysers [Johnson and Johnson Vitros Ektachem machine, Beckman LX20 analysers, Roche P800 modular system]) were used to measure serum levels of total cholesterol, high density lipoprotein (HDL) cholesterol and triglycerides (TG) on fasted blood samples using enzymatic colorimetric assays. Using an enzymatic colorimetric slide assay fasting blood glucose was measured on an Ektachem 700 multichannel analyser (Johnson and Johnson Clinical Diagnostic Systems, Amersham, UK).

3.1.6 Metabolomics

3.1.6.1 Non-targeted metabolomics

Metabolites were detected and quantified in fasted serum and plasma samples by the metabolomics platform Metabolon, Inc. (Durham, NC), which uses a non-targeted mass spectrometry based-approach as has been previously described (Menni et al., 2013b) and described in greater detail below. Metabolomics was completed cross-sectionally in three batches, one in serum and two in plasma. Quality control of the non-targeted metabolomics dataset was undertaken by Metabolon Inc. and Dr Cristina Menni. Raw data were median-normalised by dividing
metabolite concentrations by the day median of that metabolite. As the metabolite concentrations were not normally distributed, all metabolites underwent a rank-based inverse normal transformation (INT). The rank-based INT method used back transformation of the sample quantile to estimate the expected normal scores (Beasley et al., 2009). Overall there were 456 metabolites, including 292 chemically identified ('known') and 164 for which the identity is currently unknown ('unknown'). In 2016 non-targeted metabolomics profiling was also completed on blood samples at 3 time points on 2069 twins, the same quality control procedures were undertaken, the details of this dataset are not included as only one metabolite was used for analysis in Chapter 6.

3.1.6.1.1 Metabolon metabolomics detailed methods

Methanol was added to samples for protein precipitation and isolation of a range of metabolites, samples were shaken vigorously for 2 minutes, centrifuged, and the resultant extract divided into four portions: one to be analysed by ultra-high performance liquid chromatography-tandem mass spectrometry (UPLC-MS/MS; positive mode), one to be analysed by UPLC-MS/MS (negative mode), one to be analysed by gas chromatography–mass spectrometry (GC-MS), and one sample saved in case of incident. Three different controls were used in conjunction with sample analysis, these included: human plasma samples that have been analysed considerably by Metabolon, Inc. for replication; water samples for process blanks; and multiple standard solutions spiked into all experimental samples to assess equipment performance. Randomisation of samples and controls was ensured across runs.

The UPLC-MS/MS platform used a Waters Acquity UPLC and a ThermoFisher LTQ mass spectrometer, the latter comprised of an electrospray ionization source and a linear ion-trap mass analyser. The instrument had scanning set at 99-1000 m/z and to fluctuate between MS and MS/MS. Positive and negative ions were identified in acidic and basic extracts, respectively by separate injections. Columns (Waters UPLC BEH C18-2.1 × 100 mm, 1.7 µm) had sample extracts loaded and then gradient-eluted with water and 95% methanol with 0.1% formic acid or 6.5 mM ammonium bicarbonate for acidic and basic extracts, respectively. Following each injection, columns were washed.

Analysis of samples by GC-MS were first vacuum dried for 18 h minimum before derivitization using bistimethyl-silyl trifluoroacetamide under nitrogen. Following derivitization,

separation of samples occurred on a 5% phenyldimethyl silicone column. The column used helium as the carrier gas at a temperature rise from 60° to 340° C for a run time of 17 minutes. Sample analysis was conducted on a Thermo-Finnigan Trace DSQ MS set at unit mass resolving power with electron impact ionization and an atomic mass unit scan range from 50-750.

Software developed at Metabolon was used for metabolite identification, which compared ion features (retention time, molecular weight (m/z), preferred adducts, MS spectra, etc.) in the experimental samples to a reference library (Dehaven et al., 2010). Known metabolites were identified by comparison to a mass spectroscopy library containing >2,400 purified standards. Approximately 5,300 unknown, but commonly reoccurring biochemicals have been added to this library for identification in the experimental samples. Peaks were quantified by area under the curve and therefore use relative measures and hence have no units.

3.1.6.2 Targeted metabolomics

A targeted metabolomic assay using the Biocrates Absolute IDQTM-kit p150 (BIOCRATES Life Sciences, AG, Innsbruck, Austria) was conducted on serum samples (by Biocrates employees) from 1030 twins who also had non-targeted metabolomics profiling from the same visit the details have been previously described (Illig et al., 2010, Römisch-Margl et al., 2012). In short, the flow injection analysis (FIA) tandem mass spectrometry (MS/MS) method quantifies 163 small molecule metabolites at once through multiple reaction monitoring. Metabolite quantification is completed by reference to the applicable internal standards. Concentrations of targeted metabolites are reported in µM.

The Biocrates platform analyses acylcarnitines (Cx:y), hydroxylacylcarnitines [C(OH)x:y] and dicarboxylacylcarnitines (Cx:y-DC); amino acids; sphingomyelins (SMx:y) and sphingomyelinderivatives [SM(OH)x:y]; and glycerophospholipids (PC) in absolute metabolite values (mM). The metabolite concentrations were right-skewed and therefore log transformation was undertaken. There were 18 metabolites measured both by the Biocrates and Metabolon platforms, therefore for analyses including both platforms only 145 metabolites were considered from the Biocrates platform. **Table 3-1** shows the Pearson's correlation between metabolites measured on both platforms.

Metabolite name	Pearson correlation (P<0.001)
Glutamine	0.3121
Tryptophan	0.5094
Histidine	0.3099
Phenylalanine	0.4804
Threonine	0.5537
Tyrosine	0.6411
Methionine	0.5298
Ornithine	0.1522
Valine	0.6377
Proline	0.8113
Acetylcarnitine	0.6423
Serine	0.4613
Hexanoylcarnitine	0.6972
Glycine	0.6759
Butyrylcarnitine	0.6375
Propionylcarnitine	0.6719
Octanoylcarnitine	0.7703
Decanoylcarnitine	0.7743

Table 3-1. Pearson's correlation between metabolites measured by the targeted and nontargeted platforms (*n*=1030)

3.1.7 Genotyping

Genotyping for TwinsUK samples has been outlined recently (Metrustry et al., 2014). Quality control of the genotyping data was undertaken by Dr Massimo Mangino. Multiple Illumina arrays were used for genotyping: Human-Hap300 (Richards et al., 2008), HumanHap610Q, 1M-Duo and 1.2M-Duo 1M. First, normalized intensity data from each of the arrays were pooled independently (1M-Duo and 1.2M-Duo 1 M pooled together), then genotypes assigned for each dataset by the Illuminus calling algorithm (Teo et al., 2007). Had an individual's expected genotype been called with lower than a posterior probability cutoff of 0.95, no calls were assigned. Pooling was validated by a visual inspection for observable batch effects of 100 random, shared SNPs. SNPs were excluded in a similar manner from each of the three datasets by visually inspecting intensity cluster plots of significant SNPs for no calling bias due to overdispersion, and/or false assignment of genotype. Samples and SNPs were further excluded for the reasons outlined in **Table 3-2** (columns **A** and **B**, respectively). Each of the three datasets had alleles aligned to HapMap2 or HapMap3 forward strand alleles. A pair-wise comparison was then conducted between the three datasets that allowed

for additional sample and SNP exclusion for subsequent avoidance of false genotyping, as outlined in **Table 3-2** (column **C**). Finally, the three datasets were merged, reserving data from the array that had typed the largest number of SNPs when two different arrays were used for one individual.

Table 3-2. Exclusion criteria for samples, SNPs and following pair-wise comparison

A. Samples	B. SNPs	C. Pair-wise comparison			
1. Sample call rate <98%	 Hardy-Weinberg p-value <10⁻⁶ (determined in a sample of unrelated subjects) 	1. Similarity at corresponding samples <1%			
 2. Heterozygosity across all SNPs ≥ 2 SD from the sample mean 	2. MAF <1% (determined in a sample of unrelated subjects)	2. Similarity at corresponding SNPs <1%			
3. Identification of non-European ancestry by comparison of the PCA with HapMap3 populations.	3. SNP call rate < 97% for SNPs with MAF \ge 5%, or <99% for 1% \le MAF < 5%	3. All pair-wise dataset comparisons inspected for logistic regression by visual investigation of QQ plots			
4. Pair-wise IBD probabilities imply sample misidentification		4. Hardy-Weinberg p-value <10 ⁻⁶ (determined in a sample of unrelated subjects)			
5. Misidentified MZ and DZ twins were amended in accordance with IBD probabilities		5. Pair-wise IBD probabilities imply sample misidentification			

Abbreviations: SNPs, single nucleotide polymorphisms; SD, standard deviation; PCA, Principal Components Analysis; IBD, identity-by-descendant; MZ, monozygotic; DZ, dizygotic; MAF, minor allele frequency; Quintile-Quintile, QQ.

3.2 Statistical analysis

3.2.1 Heritability

Heritabilities of dietary phenotypes were determined using linear structural equation modeling in Mx (Neale et al., 2003, Neale et al., 1992). The variance of the phenotype is decomposed into 3 different effect components: A, additive genetic; C, common environmental; E, non-shared environmental. This is termed univariate ACE modelling. If MZ twins are significantly more alike in a trait than DZ twins, it is indicated that additive genetic effects are high. The common environmental component indicates the effect of the family environment on a trait. In the model the effect of the family is assumed to be equal in both MZ and DZ twin pairs (Kyvik, 2000). The effects that are unique to each individual and also the measurement error are captured by the non-shared environmental component. Heritability (h^2) is the proportion of the variance in a trait due to genetic factors and is demonstrated by the equation, $h^2 = (A)/(A + C + E)$. Each of the four models: ACE, AE, CE, and E were tested and evaluated by the Akaike's information criterion (AIC); the lowest AIC

indicated the best fitting model reflecting a good balance between goodness of fit and parsimony (Neale et al., 1992). Prior to modelling, the phenotypes were adjusted for age and sex using linear regression and the residuals used in the subsequent analysis.

Chapter 4 Metabolomic associations of self reported food group intakes and food patterns

In this chapter I have undertaken a discovery approach, examining all metabolite associations with food intakes and dietary patterns in the TwinsUK population with the primary aim of identifying candidate food intake biomarkers. I applied the discordant MZ twin model as a novel method of replicating significant findings from the discovery analysis.

Part of this work has been published in PLoS One (Pallister et al., 2016).

4.1 Introduction

For decades epidemiological studies have depended on self-reported accounts of food intake to evaluate dietary intakes. Using these approaches, however it is difficult to precisely capture food intakes and portion sizes and consequently the findings of diet-disease association studies have been called into question. Using objective biomarkers of food intake (such as essential fatty acids in blood to estimate fatty fish intake), there is potential to improve upon this issue, but not many biomarkers have so far been identified and have scarcely been utilized in epidemiological settings.

The high-throughput profiling of metabolites completed through targeted and non-targeted means in tissues and biological fluids used in collaboration with dietary studies is deemed nutritional metabolomics. Diet and the chemicals within our bodies (metabolites) are intimately linked, as the food we consume provides the basic chemical inputs which our bodies use to produce metabolic products downstream, in addition to the energy required to complete these mechanisms that are essential to life.

Recently, metabolomics studies using non-targeted approaches have identified novel biomarkers of food intake in large population settings. Among those, subjects from the Prostate, Lung, Colorectal, and Ovarian Cancer Screening Trial were used to identify 39 candidate dietary biomarkers for multiple food groups (Guertin et al., 2014b). In two separate studies of subjects from the African Americans in the Atherosclerosis Risk in Communities Study 39 metabolites were found to associate with alcohol intake (Zheng et al., 2014) and 48 to reported food intakes (Zheng et al.,

2014). Studies that have applied targeted metabolomics methods have also been successful in identifying significant associations between self-reported dietary intake patterns and serum metabolites (O'Sullivan et al., 2011, Altmaier et al., 2011, Floegel et al., 2013b).

A primary issue to be addressed is how candidate biomarkers of a healthy diet can be most effectively identified in collaboration with currently utilised methods of dietary assessment. Emerging research on dietary patterns is showing promising results within classic nutritional epidemiological studies (Hu, 2002). Empirically-derived dietary patterns are particularly relevant as they represent real trends in population reporting. Conversely, an advantage of using *a priori* over data-driven methods to detect metabolite biomarkers is that these patterns can be replicated in other populations. Due to its emerging importance for ameliorating derangements in metabolism associated with the metabolic syndrome (Kastorini et al., 2011), and the tendency of subjects in this group to underreport intakes (Kastorini et al., 2011), objective biomarkers of the Mediterranean diet would be particularly useful for clinicians and nutritionists.

Genetic factors have a strong impact on metabolism and may influence up to 81% of the variation in blood levels (Shin et al., 2014). As shown by classic examples of gene mutations causing inborn errors of metabolism that need dietary adjustments (e.g. phenylketonuria, maple syrup urine disease), complicated interactions exist between genes, metabolism and diet. However, recent studies suggest variation at a multiple loci with less exaggerated independent effects on metabolism are likely primarily responsible for the interplay between diet and its link to complex diseases (Kettunen et al., 2012). The effect of lipid metabolism genes (e.g. hepatic lipase gene, cholesteryl ester transfer protein) on the variation in blood cholesterol levels has been shown to depend on dietary fat content in recent intervention studies (Qi et al., 2015, Xu et al., 2015), however effects were minimal. Using more than 400 blood metabolites in the TwinsUK and the Cooperative health research in the Region of Augsburg (KORA) datasets (Wichmann et al., 2005, Shin et al., 2014), 145 metabolic loci were shown to associate with blood metabolites. The identified loci from these studies may pinpoint areas where metabolism, diet and genetics interact.

It may be challenging to replicate findings between populations due to the large interindividual variability in metabolite levels (Sampson et al., 2013), to factors such as age (Menni et al., 2013b) and genetics (Shin et al., 2014). Identical (monozygotic, MZ) twins who are matched for age, sex and the baseline genetic sequence, may aid reproducibility issues by acting as "co-twin

controls". The TwinsUK group has used this method previously in one dietary metabolomics study (Menni et al., 2013c), where FFQ-derived principal component dietary patterns were found to correlated with 42 metabolites in the greater twin population with top findings replicated in MZ twins discordant for the dietary patterns.

Using blood samples profiled by both targeted and non-targeted metabolomic platforms, my aims for this chapter were to:

- To identify novel associations with blood metabolites with reported food intakes and dietary patterns.
- ii) To replicate these associations using MZ twins discordant for food intakes.
- iii) To identify potential genotypes influencing food intakes from SNPs previously shown to be associated with metabolite levels on the TwinsUK and KORA cohorts (Shin et al., 2014).
- iv) To provide the results of this analysis online through designing the DietMetab tool.

4.2 Materials and methods

I included 3559 female twins, who completed a 131-item validated FFQ between 1995 and 2007, and had Metabolon metabolomics and BMI data available within and including (+/-) 5 years of FFQ completion. I also analysed a subset of this sample who had Biocrates targeted metabolomics data (n=858).

The characteristics of the study population can be found in **Appendix C Table 1**.

4.2.1 Food items

Previously, intake frequencies were adjusted for total energy intake using the residual method (Willett and Stampfer, 1986) and I summed these into 71 food groups based on nutrient content, taste and usage (**Appendix C Table 2**).

4.2.2 Dietary patterns

4.2.2.1 Principal components

The principal components analysis (PCA) had been completed and the steps outlined previously (Teucher et al., 2007). The PCA was undertaken by nutritionists at the University of East Anglia. Briefly, the 131 food items were combined a priori into 54 groups based on nutrient content and culinary use. Intake frequencies of these groups were the sum of the servings per week for the original questionnaire items they were derived from. The residual energy adjusted food group intake frequencies (servings/week) were standardised to z scores and then used in the PCA. Bootstrapping is a resampling approach used to approximate properties of an estimator by evaluating those properties when sampling from an approximating distribution. It was used to test if family relatedness influenced the results. Briefly, the sampling distribution of the components was estimated in unrelated individuals. 10,000 bootstrap replications of the PCA were undertaken on subsamples containing one randomly selected twin. The full dataset PCA loadings were assessed against a 95% confidence interval determined from the 2.5% and 97.5% percentiles of this distribution. It was determined that family relatedness had no significant effect. The top 20 components were examined on a scree plot (Appendix C Figure 1), elbows were shown after components 1 and 5, however, components 2 to 5 were reasonable dietary patterns therefore the first 5 components were used. These 5 components, that explained 22% of the total variance, were denoted according to their top factor loadings

(Table 4-1) as follows: 'Fruit and vegetable'; 'High alcohol'; 'Traditional English'; 'Dieting'; and

'Low meat'. Consult **Appendix C Table 3** for their complete factor loadings.

Diet	Variance		
pattern	explained	High intakes ¹	Low intakes ²
Fruit & vegetable	8.2%	Fruit, allium and cruciferous vegetables	Fried potatoes
High alcohol	3.9%	Beer, wine and allium vegetables	High fiber breakfast cereals and fruit
Traditional English	3.6%	Fried fish and potatoes, meats, savoury pies and cruciferous vegetables	
Dieting	3.3%	Low-fat dairy products, low- sugar soda	Butter and sweet baked products
Low meat	3.2%	Baked beans, pizza and soy foods	Meat, other fish and seafood, and poultry

 Table 4-1. Principal component derived dietary patterns, with percentage of variance explained, and foods consumed in high and low intakes (Teucher et al., 2007)

¹Food frequency questionnaire items with factor loadings ≥ 0.20 .

²Food frequency questionnaire items with factor loadings \leq -0.20.

4.2.2.2 Mediterranean diet score

I calculated Mediterranean diet adherence using the modified Mediterranean diet score (MDS) method, as outlined by Trichopoulou (Trichopoulou et al., 2005). Nine food/nutrient categories were included to generate a Mediterranean diet score (MDS) of 0 to 9 ("least" to "most" Mediterranean). The two nutrient intake categories included alcohol and fatty acid ratio (monounsaturated+polyunsaturated/saturated fatty acid intakes), both in grams per day and residual energy adjusted. Seven food group intakes were generated including: fruits (including nuts), vegetables, meats, fish, dairy, cereals, and legumes. To form groups, I first multiplied intake frequencies for assigned foods by the amount in grams per day (raw data are per week). Next, the amount in grams of these foods was added to make the final category total. Categories in grams were then adjusted using the residual method (consult Section 3.1.2.2).

I then generated the scores. For all frequency categories as well as the fatty acid intake ratio, I determined median intakes of each category using all eligible twin data for FFQs 1 and 2 combined. I assigned a score of 0 (no MDS) or 1 (MDS) for each category depending on whether the twin was above or below the median intake, specific to the category (refer to **Table 4-2**). For alcohol intakes, a range was used for score assignment: twins between 5 and 25 g/d were assigned a score of 1, while those above or below this range were assigned a score of 0.

Finally, the MDS was then generated by summation of each category score to generate a score

of 0 to 9.

U		
Above median	Below median	Moderate
Fruit & nuts	Meat	Alcohol: 5 – 25 grams/d
Vegetables	Dairy	-
Legumes		
Fish		
Cereals		
Unsaturated : saturated fatty acid		

Table 4-2. Criteria for a score assignment of 1 on the Mediterranean diet

4.2.3 Replication population

One metabolite was particularly strongly associated with low fat milk intake in the TwinsUK sample, I therefore obtained data on milk intake from the Cooperative Health Research in the Region of Augsburg (KORA) study to replicate this association. The KORA study includes individuals from the region of Augsburg, Germany who are unrelated (Holle et al., 2005). As part of the fourth survey (S4) between 1999-2001 4261 persons aged 25-74 years were examined. For the replication analysis, 1593 individuals with serum metabolite levels of the top milk-associated metabolite in the TwinsUK sample were used. Dietary intake in the KORA study was determined by a validated FFQ on the same day as blood sampling (Winkler and Doring, 1998)

The questionnaire included 24 food items and asked subjects to recall their "average intake" out of the six frequency categories: almost daily, several times per week, about once a week, several times per month, once a month or less, never. For the replication analysis I used the item "milk intake including buttermilk". Prior to analysis, I recoded the milk intake variable so interpretation of the result was easier therefore the least frequency ("never") was coded as 1 and the highest frequency ("almost daily") was coded as 6, and I changed the rest of the categories as follows: once a month or less, 2; several times per month, 3; about once a week, 4; and several times per week, 5.

4.2.4 Statistical analysis

Statistical analysis was carried out using Stata version 12.

For each metabolite, I ran a random intercept linear regression analysis using each food group as the predictor in the discovery sample, excluding discordant MZ twin pairs (pairs \geq 1 SD apart in intake by food group). I adjusted for age, metabolite batch effects, BMI and family relatedness: $Y_i = \beta_0 + \beta_i X_{ij} + \gamma_i age_{ij} + \delta_i BMI_{ij} + \zeta_j + \varepsilon_{ij}$

where Y_i is the metabolite and X_{ij} is the food group intake of twin *j* from pair *i*, and ζ_j is the family-specific error component that captures the unobserved heterogeneity or family characteristics.

I used the Bonferroni correction to adjust for multiple testing which allowed a significant threshold of 1.08×10^{-6} (0.05/[77 diet phenotypes x 601 detected metabolites]). For associations that passed the significance threshold in the discovery sample, I repeated/replicated them using the same model in the MZ discordant twins sample. Associations were considered replicated in the MZ discordant sample if they (i) passed the 5% level of significance threshold and (ii) had the same effect direction as the discovery group (I used only this second criteria for the targeted platform). Lastly, using an inverse variance fixed effects meta-analysis I combined the results of both analyses, these are the results that I report here. To test if dietary patterns associated to a unique metabolomic signature not captured by investigating single food groups I repeated the analysis for each of the six diet patterns (five principal components and the MDS). The beta coefficients (β) I present for the significant associations represent the reported food amount in servings per week or dietary pattern score that corresponds to a 1 SD change in the metabolite level.

4.2.5 Diet pattern association with metabolite mediated by reported food intake

For each metabolite associated with both diet patterns and food group intakes first the proportion of the variance of metabolite was determined for each relevant diet pattern after taking into account all covariates (age, BMI, batch effects, family relatedness). This quantity is indicated as r_x^2 . The proportion of the variance for the metabolite explained by the applicable diet pattern was then calculated after taking into account the same covariates as above but also including intakes of all associated foods (r_x^2) . The percentage of the diet pattern association mediated by the food group intake (r_y^2) was calculated as the proportion of the variance of metabolite that is due to the diet pattern association with the food intake, namely $1 - (r_{xy}^2/r_x^2)$.

4.2.6 Genotype associations

I have outlined the genotyping protocols that have been undertaken in the TwinsUK population in **Section 3.1.7**. A genome-wide association study (GWAS) was previously undertaken on the Metabolon metabolomics datasets conducted for TwinsUK and the Cooperative health research in the Region of Augsburg (KORA) cohorts (Wichmann et al., 2005, Shin et al., 2014). Based on the results of this study, I undertook diet-genotype associations on gene variants (56 SNPs) that were associated with blood levels of dietary-associated metabolites (48 metabolites). To identify associations, I used additive genotype as a predictor of the appropriate energy-adjusted food group intake or dietary pattern, including age and family relatedness as covariates. I defined statistical significance using a Bonferroni correction of 2.94×10^{-4} (0.05/170 tests).

4.3 Results

4.3.1 Food group-metabolite associations

I found 178 significant associations with 39 food groups after meta-analysing the discovery and MZ discordant twin groups, consisting of 105 unique metabolites (**Appendix C Tables 5, 6, and 7**). From these 105 unique metabolites, 73 have been identified (**Figure 4-1**) and 32 are unknown at this time. Those 73 identified metabolites belong to six biochemical groups: 38 lipids, 16 amino acids, 14 xenobiotics, 3 carbohydrates, 1 cofactor/vitamin, and 1 peptide (**Figure 4-2a and b**). To my knowledge, 72 of the food associations with identified metabolites have not been previously identified in nutritional metabolomics using similar methods (**Appendix C Table 7**).





by the metabolic role of the metabolite.

I found reported intakes of alcoholic beverages to be associated with the most metabolites, including 41 associations overall (38 wine,1 spirits/liquors) with 15 associations (of the known metabolites) I believe to be novel; the top association was between wine and scylloinositol (meta-analysis result: Beta[SE]: 0.052[0.003]; P=1.47x10⁻⁴⁹). Reported intakes of teas and coffee included 27 associations (8 novel of the known metabolites) overall including: 12 coffee, 9 black tea, and 6 herbal tea. Notably, for both coffee and tea, 6 metabolites belonging to the xenobiotics pathway, i.e. metabolites foreign to the human body, were identified. These likely originated from gut bacterial metabolism of caffeine (e.g. catechol sulfate, 1methylxanthine). The top association was between coffee and the unknown metabolite X-14473 (0.038[0.001]; P=6.12x10⁻¹⁸⁷). There were 26 associations (4 novel) with reported seafood consumption, including: 15 oily fish and 11 with other seafood. A number of the metabolites associated with reported seafood consumption were essential fatty acid metabolites; the top association was between docosahexaenoate (DHA; 22:6n3) and oily fish (0.177[0.013]; $P=2.09 \times 10^{-44}$). I found 8 associations (5 novel) with meat products, specifically: 3 meat, 2 poultry, 2 processed meat and 1 beef burgers. The associations were primarily with amino acids, the top association being a novel associations between meat and trans-4-hydroxyproline (0.075[0.009]; P=1.08x10⁻¹⁷). I found 14 metabolites (5 novel) were associated with reported dairy product intake, including: 9 butter, 3 cream and 2 low-fat milk. Primarily associations with dairy products included lipids, the top association was reported between low fat milk intake and the recently named metabolite trimethyl-N-aminovalerate (X-21365: 0.076[0.007]; P=9.36x10⁻ ²⁷). There were 10 associations (3 novel) with grain-rich foods, including: 5 high fibre breakfast cereals, 2 refined bread and grains, 2 porridge and 1 wholemeal bread and grains. The top association was between porridge and the unknown metabolite X-09789 (0.094[0.008]; $P=4.96 \times 10^{-33}$). There were 13 associations (4 novel) with reported fruit intakes consumption, including: 6 apple and pears, 3 citrus fruits, 1 bananas, 1 berries, 1 peaches and 1 fruit juices. The top association within the fruit category was between fruit juice and stachydrine (0.058[0.005]; P=3.26x10⁻³⁷). I identified 16 associations (8 novel) with reported vegetable intakes, including: 6 green leafy, 5 avocado, 3 allium, 1 tomatoes and 1 mushrooms. The strongest association within the vegetables group was between reporeted mushroom consumption and ergothioneine (0.181 [0.019]; P=5.93x10⁻²²). I found 17 associations (9 novel) with reported sweet and savoury discretionary food intakes, including: 5 sweet baked products, 4 savory pies, 4 fried fish, 3 confectionary and jam, and 1 savoury snacks. Within the

discretionary foods group, the strongest association was between reported savoury snack intake and the unknown metabolite X-11372 (0.051[0.007]; $P=3.88\times10^{-14}$). Other notable associations included 2 metabolites with reported chocolate intake (1 novel; top association with theobromine: 0.024[0.003]; $P=1.34\times10^{-11}$), a novel association between reported soymilk intake and 4-ethylphenylsulfate (0.239[0.033]; $P=6.05\times10^{-13}$) and also an association between an unknown metabolite and reported intake of soyfoods (X-11381: -0.108[0.020]; $P=5.80\times10^{-8}$) and nuts (X-11315: 0.054[0.005]; $P=3.75\times10^{-25}$).

A number of metabolites were associated to more than one food item, many of these are listed in **Tables 4-3** and **4-4**. Notably, the essential fatty acids eicosapentaenoate (EPA) and docosahexaenoate (DHA), their downstream metabolite 1-docosahexaenoylglycerophosphocholine and 3-carboxy-4-methyl-5-propyl-2-furanpropanoate were all positively associated with oily fish and other fish/seafood intakes, and all but DHA with avocado. These multiple associations may be due to correlated food intakes, though each source may be contributing independently. Higher levels of EPA and DHA were also associated with lower intakes of sweet baked products. The metabolite 3-phenylpropionate was associated positively to intakes of apples/pears and negatively with fried fish and savoury pies. The latter associations may be due to systematic under reporting rather than biological in origin as is discussed later. A further example of this observation are the metabolites quinate, cyclo(leu-pro), 3-methyl catechol sulfate 1 and 3-hydroxypyridine sulfate, which were positively associated with coffee intake and negatively with tea intake.

4.3.1.1 Milk association replication

I replicated the association between the previously unknown metabolite trimethyl-Naminovalerate (X-21365) and milk intake in the KORA population (0.008[0.002]; P=6.88x10⁻⁶), establishing the quality of the data.

4.3.2 Dietary pattern results

4.3.2.1 Nutrient profile of the Mediterranean diet

Energy-adjusted nutrient intakes are represented as percentages of the recommended intakes by tertile of the Mediterranean Diet Score (MDS) are presented in **Figure 4-3** with significance level for trend. Nutrient profiles of the FFQ PCs have been described previously (Teucher et al., 2007). There were significant trends for most nutrients from the bottom to top tertile of the MDS,

suggesting that individuals who scored higher on the MDS reported consuming more nutrientdense diets, high in fibre and lower in total, saturated fat and sucrose content. Despite the monounsaturated fat content being lower in those who scored in the top versus bottom tertile of the MDS, an important characteristic of the Mediterranean diet derived from olive oil intake, a logistic regression analysis of olive oil intake (yes or no) was strongly predicted by the MDS (OR= 1.15, SE= 0.02, P=1.19x10⁻¹³).

4.3.2.2 Heritability of the MDS

The best fitting model for MDS was the AE model, with heritability estimates of 0.41[95% CI = 0.36; 0.46] and 0.59[0.54;0.64] respectively.

4.3.2.1 Dietary pattern-metabolite associations

I identified 108 significant dietary pattern-metabolite associations that passed the Bonferroni level of significance ($P < 1.08 \times 10^{-6}$) in the discovery twin population and following meta-analysis (Appendix C Tables 8, 9). The MDS (Figure 4-4a) was associated with 18 metabolites (2 amino acids, 5 lipids, 2 xenobiotics and 9 unknown metabolites). Of the known metabolites, MDS was most strongly associated with DHA (meta-analysis result: 0.097[0.009]; P=2.45x10⁻ ²⁷). The F&V diet pattern (**Figure 4-4b**) was associated with 33 metabolites (6 amino acids, 3 carbohydrates, 1 cofactor/vitamin, 7 lipids, 1 peptides, 3 xenobiotics and 12 unnamed metabolites), a number of which were also associated with the MDS. A high F&V pattern was most strongly associated with glycerate (0.094[0.009]; P=1.97x10⁻²⁷), a carbohydrate involved in glycolysis, gluconeogenesis, and pyruvate metabolism. The high alcohol pattern (Figure 4-4c) was associated with 23 metabolites (3 amino acids, 1 cofactor/vitamin, 4 lipids, 3 xenobiotics, and 12 unnamed metabolites). The strongest known metabolite association with a high alcohol diet pattern was with the amino acid involved in valine, leucine and isoleucine metabolism, αhydroxyisovalerate (0.158[0.013]; P=3.20x10⁻³⁵). The traditional English diet pattern (Figure 4-4d) was associated with 6 metabolites (2 amino acids, 1 xenobiotic, and 3 unnamed metabolites). A negative association was the strongest observed between the traditional English pattern and stachydrine (-0.094[0.012]; P=9.94x10⁻¹⁶), an amino acid involved in urea cycle and arginine, proline metabolism. The dieting pattern (Figure 4-4e) was associated with 5 metabolites (2 xenobiotics, and 3 unnamed metabolites). Dieting was most strongly, negatively associated with quinate (-0.095[0.015]; P=1.40x10⁻¹⁰), a xenobiotic derived from benzoate metabolism. The low meat pattern (Figure 4-4f) was associated with 22 metabolites (7 amino

acids, 7 lipids, and 8 unnamed metabolites). Most prominent, was a strong negative association with the furan fatty acid 3-carboxy-4-methyl-5-propyl-2-furanpropanoate (CMPF) (-0.144[0.015] P=3.63x10⁻²¹).





Associations are significant at the Bonferroni level: 1.08×10^{-6} for discovery and meta-analysis. Significance level for discordant twin replication set (*P*<0.05). Names are colour coded according to direction of association: red, positive; black, negative.

4.3.2.2 Metabolites uniquely associated with food intakes or dietary patterns and those associated to both

Overall 126 metabolites (43 unknown) were associated to at least one dietary phenotype (**Figure 4-5**). Fifty-five metabolites were associated with single food intakes but not the dietary patterns (**Figure 4-5a**). This may suggest these metabolites are less important for dietary intakes of the whole population. Twenty-one metabolites were not associated to single foods (**Figure 4-5b**) but were associated with the dietary patterns. In this case the combined food intakes may have an effect on these metabolites not occurring by the food intake alone or the associations may originate from food intakes not explicitly measured by the FFQ or behaviours correlated to a dietary pattern. Fifty metabolites were associated to at least one pattern and one food (**Figure 4-5c**), there are likely the most important metabolites related to food intake in our dataset.

4.3.2.3 Dietary pattern-metabolite associations mediated by food intakes

To confirm if the 50 metabolites associated with dietary patterns and at least one food group were driven by the reported food intakes and not other factors, I analysed the degree to which the variance in dietary pattern-metabolite associations was mediated by the metabolites association with single food intakes. I found that between <0% and 100% of the variance in the association between the metabolites and the dietary pattern scores were mediated by the metabolites association between the metabolites and the dietary pattern scores were mediated by the metabolites association with food intake (**Tables 4-3** and **4-4**). The MDS and fruit and vegetable diet pattern associations with essential fatty acid metabolites (eicosapentaenoate, 1-docosahexaenoylglycerophosphocholine*, CMPF and docosahexaenoate) were strongly accounted for by food intakes (range: 86.2% to 99.7%). This was also the case for the low meat pattern association with pyroglutamine* which was strongly accounted for by meat intakes (93.2%), the dieting pattern with quinate which was accounted for by coffee and tea intakes (97.9%), the high alcohol pattern with 4-androsten-3beta, 17beta-diol disulfate 1*, 5alpha-androstan-3beta, 17beta-diol disulfate and scyllo-inositol accounted for by wine or spirit intake (98.3%, 92.4% and 99.5%, respectively).

4.3.2.4 Gene-variants SNP associations

A GWAS study was previously performed on the TwinsUK dataset to identify SNPs associated with blood metabolites levels from the Metabolon platform (Shin et al., 2014). Of these 126

dietary-associated metabolites, 48 contributed to 62 metabolite-SNP associations including 56 unique SNPs in 39 genes (**Appendix C Table 10**). I undertook this analysis to identify potential SNPs which may be influencing dietary intakes. I did not find any SNP-diet associations with these variants at the Bonferroni level of statistical significance (P<2.94x10⁻⁴ = 0.05/170 tests). I found 19 SNP-diet associations at the nominal level (P<0.05; **Appendix C Table 11**). The fact that I did not identify any significant associations between diet and metabolite-related SNPs suggests that these SNPs do not have such a strong effect as to influence eating behaviour, although it could also be due to the limited power due to the low sample size or the lack of precision by FFQs.



Table 4-3. List of known metabolites associated with food intakes and dietary patterns and the degree to which their association with each pattern is through reported food intake

Metabolite	No.	Foods associated	R ² all	Diet patterns associated	R ² pattern no foods (1)	R ² pattern with foods ⁽²⁾	% association
eicosapentaenoate	6	↑ Avocado, High fat salad	0.08	↑ MDS	0.026	0.001	95.1
		dressing, Oily fish, Other fish/seafood, Wine		↑ F & V	0.047	0.004	90.9
		↓ Sweet baked products		↓ Low Meat	0.025	0.004	86.1
1-	4	↑ Avocado, Green leafy	0.04	↑ F & V	0.023	0.002	93.2
docosahexaenoylgly cerophospho- choline*		vegetables, Olly fish, Other fish/seafood		\uparrow MDS	0.022	0.003	86.2
3-carboxy-4-methyl- 5-propyl-2-	4	↑ Avocado, Green leafy vegetables, Oilv fish, Other	0.10	↑ F & V	0.029	0.000	99.7
furanpropanoate		fish/seafood		↑ MDS	0.028	0.001	97.5
				↓ Low Meat	0.036	0.015	59.2
docosahexaenoate (DHA: 22:6n3)	4	↑ Oily fish, Other fish/seafood, Wine	0.11	↑ F & V	0.043	0.003	93.4
())		↓ Sweet baked products		↑ MDS	0.035	0.003	91.1
				↓ Low Meat	0.027	0.004	85.0
3-phenylpropionate (hydrocinnamate)	3	↑ Apples/pears ↓ Fried fish, Savoury pies	0.03	↑ F & V	0.027	0.006	76.8
				↓ Traditional English	0.015	0.006	62.0
glycerate	3	↑ Citrus fruit, Tomatoes	0.03	↑ F & V	0.037	0.009	75.8
pyroglutamine*	3	↓ Meat, Other fish/seafood,	0.04	↑ Low Meat	0.022	0.002	93.2
scyllo-inositol	3	Poulty ↑ Wine	0.13	↑ High	0.043	0.000	99.5
		↓ Fried fish, Sweet baked products		Alconol ↑ F & V	0.035	0.010	71.3
				↑ MDS	0.015	0.010	30.6
4-androsten- 3beta,17beta-diol disulfate 1*	2	↑ Spirits/liquor, Wine	0.09	↑ High Alcohol	0.035	0.001	98.3
creatine	2	↑ Meat, Poultry	0.03	↑ Traditional English	0.014	0.001	89.9
	_			↓ Low Meat	0.015	0.002	87.9
ergothioneine	2	↑ Mushrooms ↓ Savoury pies	0.12	↑ F & V	0.048	0.006	88.4
				↑ MDS	0.025	0.007	71.1
	_		0.00	↑ High Alcohol	0.041	0.022	47.0
indolepropionate	2	↑ Apples/pears, Bananas	0.03	↑ F & V	0.029	0.012	58.2
				↑ MDS	0.021	0.011	50.5
pipecolate	2	↑ Wine ↓ Confectionary/jam,	0.03	↑ F & V	0.016	0.007	58.5
quinate	2	↑ Coffee ↓ Black tea	0.20	↓ Dieting	0.014	0.000	97.9
stachydrine	2	↑ Citrus fruit, Fruit juice	0.11	↓ Traditional English	0.019	0.010	45.5
1,5-anhydroglucitol	1	↓ Other fish/seafood	0.01	↓F&V	0.022	0.013	41.6
10-undecenoate (11:1n1)	1	↑ Butter	0.02	↑ High Alcobol	0.011	0.006	42.5
15-methylpalmitate (isobar with 2-	1	↑ Butter	0.03	↓ Low Meat	0.016	0.009	46.0
2-aminobutyrate	1	↑ Wine	0.02	↑ High	0.017	0.003	83.1
				Alcohol ↓ Low Meat	0.018	0.008	57.4
2-hydroxybutyrate	1	↑ Wine	0.02	↓ Low Meat	0.011	0.004	60.2
4-ethylphenylsulfate	1	↑ Soy/other milk	0.06	↑ Dieting	0.015	0.003	81.0
5alpha-androstan- 3beta,17beta-diol disulfate	1	↑ Wine	0.06	↑ High Alcohol	0.029	0.002	92.4
alpha- hydroxvisovalerate	1	↑ Wine	0.07	↓ Low Meat	0.014	0.001	91.5
				↑ High Alcohol	0.049	0.008	84.1

Table 4-3. List of known metabolites associated with food intakes and dietary patterns and the degree to which their association with each pattern is through reported food intake

	No.		R ² all	Diet patterns	R ² pattern no foods	R ² pattern with	% association
Metabolite	foods	Foods associated	foods	associated	(1)	foods (2)	through food
catechol sulfate	1	↑ Coffee	0.03	↑ F & V	0.014	0.015	-3.5
myo-inositol	1	↑ Wine	0.02	↑ F & V	0.012	0.008	34.7
Octenoylcarnitine	1	↑ White/brown bread & refined grains	0.03	↓ F & V	0.038	0.019	50.3
Phosphatidylcholine diacyl C36:5	1	↑ Wine	0.05	↓ Low Meat	0.062	0.033	46.6
Phosphatidylcholine diacyl C38:6	1	↑ Oily fish	0.04	↑ F & V	0.031	0.008	73.4
,				↓ Low Meat	0.036	0.027	25.8
piperine	1	↑ Wine	0.03	↑ High Alcohol	0.031	0.009	71.2
				↑ MDS	0.011	0.009	16.5
threitol	1	↑ Apples/pears	0.03	↑ F & V	0.016	0.005	71.2
tryptophan betaine	1	↑ Allium vegetables	0.03	↑ F & V	0.026	0.007	74.6
				↑ High Alcohol	0.028	0.011	62.0
				↑ MDS	0.024	0.010	57.3

F & V, fruit and vegetable pattern; MDS, Mediterranean diet score

(1) The proportion of the variance in the diet pattern explained by the metabolite after taking into account all covariates (age, BMI, batch effects and family relatedness).

(2) The proportion of the variance in the diet pattern explained by the metabolite after taking into account all covariates as in (1) and adjusting for all foods associated to the metabolite.

Table 4-4. List of unknown metabolites associated with both food intakes and dietary patterns and the degree to which their association with each pattern is mediated by reported food intakes

Metabolite	RT	Mass	No. foods	Foods associated	R ² all foods	Diet patterns associated	Pattern no foods	Pattern with foods	% association through food
X-11315	1.19	130.2	13	↑ Apples/pears, Berries,	0.13	↑ F & V	0.088	0.001	98.4
				vegetables, High fibre		↑ MDS	0.042	0.002	94.6
				breakfast cereals, Nuts, Oily fish, Peaches ↓ Confectionary/jam, Fried fish, Savoury pies, Sweet baked products, White/brown bread & refined grains		↓ Traditional English	0.019	0.004	79.3
X-02269	1.55	255.1	6	↑ Allium vegetables,	0.11	↑ F & V	0.051	0.002	96.5
				vegetables, High fibre		↑ MDS	0.039	0.002	94.8
				breakfast cereals, Oily fish, Other fish/seafood		↓ Low Meat	0.020	0.005	75.9
X-09789	2.62	153.1	5	↑ Apples/pears, High	0.09	↑ F & V	0.016	0.001	98.2
				Porridge, Wholemeal bread/grains		↓ High Alcohol	0.026	0.004	90.1
X-11372	NA	NA	5	↑ Fried fish, Savoury	0.08	↓ F & V	0.046	0.004	91.6
				↓ Apples/pears, Green		↑ Traditional	0.015	0.005	65.1
				leaty vegetables		↑ Low Meat	0.016	0.014	14.4
X-11469	3.82	239.1	5	↑ Avocado, Green leafy vegetables, High fibre	0.11	↓ Low Meat	0.021	0.001	97.2
				breakfast cereals, Oily		↑ F & V	0.056	0.002	95.7
				lish, Other lish/sealood		↑ MDS	0.040	0.002	94.8
X-11381	1.11	186.2	2	↑ Processed meats ↓ Soy foods	0.03	↑ Traditional English	0.011	0.003	76.6
						↓MDS	0.017	0.011	36.4
						↓ High Alcohol	0.012	0.013	-9.2
X-11799	1.58	226.0	2	↑ Wine ↓ Sweet baked products	0.06	↑ High Alcohol	0.030	0.001	96.0
X-12816	NA	NA	2	↑ Coffee Black tea	0.15	↑ High Alcohol	0.015	0.002	85.8
X-14374	NA	NA	2	↑ Coffee	0.09	↓ Dieting	0.010	0.000	98.0
X-14473	3.27	211.2	2	↑ Coffee	0.28	↓ Dieting	0.033	0.001	97.0
X-01911	4.26	464.1	1	↑ Wine	0.03	↑ High Alcobol	0.030	0.007	77.6
X-02249	4.03	267.2	1	↑ Butter	0.02	↑ Dieting	0.013	0.004	71.8
						↓ Low Meat	0.019	0.011	40.0
X-04495	NA	NA	1	↑ Wine	0.02	↑ High Alcohol	0.016	0.001	93.8
X-10395	9.94	156.0	1	↑ Wine	0.03	↑ High Alcohol	0.017	0.001	91.7
X-11795	NA	NA	1	↑ Wine	0.06	↑ High Alcohol	0.043	0.007	83.4
X-11858	NA	NA	1	↑ Allium vegetables	0.03	↑ F & V	0.026	0.006	77.1
X-12038	5.82	245.3	1	↑ Wine	0.01	↑ High Alcohol	0.008	0.001	88.6
X-12798	1.84	240.1	240.1 1	↑ Low fat mik	0.02	↓ High Alcohol	0.020	0.008	59.4
						↓ MDS	0.016	0.008	48.8
X-21365 ⁽³⁾	NA	NA	1	↑ Low fat mik	0.04	↓ High Alcohol	0.017	0.005	73.7
						↓ MDS	0.019	0.009	54.1

RT, retention time; F & V, fruit and vegetable pattern; MDS, Mediterranean diet score; NA, not available

(1) The proportion of the variance in the diet pattern explained by the metabolite after taking into account all covariates (age, BMI, batch effects and family relatedness).

- (2) The proportion of the variance in the diet pattern explained by the metabolite after taking into account all covariates as in (1) and adjusting for all foods associated to the metabolite.
- (3) Metabolite recently identified as trimethyl-N-aminovalerate.

4.4 Discussion

Using one of the largest datasets of its kind, I identified and replicated using MZ discordant twins 73 novel associations between metabolites and specific food groups, providing further potential intake biomarkers for future research. Moreover, I found 108 significant metabolite associations with 5 different dietary patterns. Forty-eight metabolites contributed to 62 metabolite-SNP associations in 56 unique SNPs that were identified from a previous metabolomics GWAS study (Shin et al., 2014), though I did not find these SNPs to be related to dietary intakes.

4.4.1 Food items

Alcohol consumption

To my knowledge the associations between higher reported wine intake and increased branched-chain amino acids (BCAA; valine, leucine and isoleucine and their metabolites, 3-methyl-2-oxobutyrate and 4-methyl-2-oxopentanoate) and medium-chain fatty acids (caprate and caprylate) are novel findings. BCAA, 3-methyl-2-oxobutyrate and 4-methyl-2-oxopentanoate have been found to be higher in twins with type 2 diabetes or impaired fasting glucose (3-methyl-2-oxobutyrate being the strongest predictor) previously (Menni et al., 2013a) and associated with a higher BMI (Moore et al., 2014). Under impaired mitochondrial oxidation of lipids and glucose, levels of these BCAA catabolites increase, suggesting mitochondrial dysfunction. Insulin resistance has found to occur under binge drinking (Lindtner et al., 2013) though long-term effects are not known, these associations may highlight a novel pathway mediating this effect.

I also confirmed a number of metabolite associations with reported alcohol intake from a previous metabolomics study (Zheng et al., 2014) including higher intakes of wine with increased alpha-hydroxyisovalerate, scyllo-inositol, a inositol metabolite, and steroids originating from dehydroepiandrosterone (DHEA; 5-alpha-androstan-3beta,17beta-diol disulfate, 4-androsten-3beta,17beta-diol disulfate 1, 5-alpha-androstan-3beta,17beta-diol disulfate and epiandrosterone sulfate) (Zheng et al., 2014). Interestingly, a variant in the *HAO2* gene (rs12141041) encoding long-chain L-2-hydroxy acid oxidase 2 associated to alpha-hydroxyisovalerate. The *HAO2* gene has been found to be related to blood pressure regulation in animal models (Barawkar et al., 2012). A variant in the *SLC5A11* gene (rs4787294) is associated with scyllo-inositol. The *SLC5A11* gene encodes a myo- and scyllo-inositol.

transporting sodium-dependent glucose transporter. The *SLC5A11* gene has been implicated in systemic lupus erythematosus (SLE) risk. Two SNPs the *SULT2A1* (rs2547231 and rs296396) gene which is involved in the sulfation of a number of steroids and bile acids, were associated to 4-androsten-3beta,17beta-diol disulfate 1 and 5-alpha-androstan-3beta,17beta-diol disulfate levels. Ethanol feeding in rats has recently shown significantly increased expression of *SULT2A1* in the intestines and liver (Maiti and Chen, 2015), suggesting this gene may be directly involved in modulating this association.

Seafood consumption

I identified novel associations between increased reported fish and seafood intakes with reduced pro-inflammatory lvsolipids (1-arachidonoylglycerophosphoethanolamine, 1-1eicosatrienoylglycerophosphocholine, 1-linoleoylglycerophosphoethanolamine, oleoylglycerophosphoethanolamine), originating from essential fatty acid (EFA) metabolism. Lysolipids are involved in the formation of the cellular lipid bilayer, though contribute to inflammatory processes forming free lysophosphatidylcholines when cleaved by lipoproteinassociated phospholipase A2. Lysophosphatidylcholines potentially aggravate inflammatory artherosclerotic plaques (Goncalves et al., 2012). I replicated previous associations between increased reported intakes of oily fish and other seafood with higher 3-carboxy-4-methyl-5propyl-2-furanpropanoate (CMPF), a furan fatty acid, and docosahexaenoate (DHA), an EFA (Guertin et al., 2014b, Zheng et al., 2014). Variants in FADS1 (rs174538, rs174556, rs968567 and rs174535), a gene encoding a delta-5 desaturase enzyme, were associated with EFA lysolipids and DHA (Yang et al., 2015). Recently, a meta-analysis confirmed two of these variants in FADS1 (rs174538 and rs174548) modulated gene-dietary-derived EFA associations (Smith et al., 2015).

Meat consumption

I found reported meat consumption to be associated with amino acids primarily, including creatine, trans-4-hydroxyproline and pyroglutamine. In particular, I found increased red meat intake to be associated with higher trans-4-hydroxyproline, a component of collagen previously shown to increase in blood following gelatin feeding (Ohara et al., 2007). Another novel association I identified was between increased reported red meat and poultry intakes and higher creatine levels. Red meat has characteristically high amounts of creatine and lower blood levels

have been found in vegetarians (Delanghe et al., 1989). Lower blood creatine has been associated with increased insulin sensitivity (Gall et al., 2010) and in liver steatotic versus NASH patients (Kalhan et al., 2011), higher levels have been found in individuals with dilated cardiomyopathy (Alexander et al., 2011). A variant in carbamoyl-phosphate synthase 1 (*CPS1*; rs715) was found to associate with creatine. CPS1 is the first enzyme of the urea cycle, producing ammonia from urea. Expression of *CPS1* is a candidate biomarker of NAFLD (Rodriguez-Suarez et al., 2012). Finally, higher intakes of seafood and meat correlated with lower levels of pyroglutamine, though little is known about this metabolite it is a cyclic derivative of glutamine, lower levels have previously been associated with chicken intake (Guertin et al., 2014b). A variant in *SLC6A13* (rs11613331), a gene that which encodes GAT2 a gamma-aminobutyric acid and betaine transporter, was associated with pyroglutamine. Other SNPs in *SLC6A13* have been related to renal function (Franceschini et al., 2014, Liu et al., 2011).

Dairy consumption

I identified novel associations between higher reported intakes of creams (double and clotted cream) with increased levels of lysoPhosphatidylcholine acyl C17:0 and C28:1 and hydroxysphingomyeline C14:1. A previous study on the EPIC-Potsdam cohort showed a diet pattern high in butter and high-fat dairy products and low margarine intake to be positively associated with lysoPhosphatidylcholine acyl C17:0 (Floegel et al., 2013b). Heptadecanoid acid (C17:0) has shown to be elevated following a dairy intervention trial, the lipid is formed by ruminal bacteria and therefore thought to be milk fat specific (Abdullah et al., 2015).

Reported butter consumption was uniquely associated with elevated levels of six lipids, including associations with increased butter intake and nonanoylcarnitine, an ester of carnitine with pelargonic acid (C9), 10-nonadecenoate (19:1n9), a monounsaturate of nonadecenoate (19:0), and myristate (14:0), a saturated fatty acid found most animal and vegetable fats. A variant in *ACADL* (rs3738934), a gene with important roles in lipid oxidation was associated with blood nonanoylcarnitine. Expression of *ACADL* has shown to be lower in mouse liver and adipocytes following high fat feeding, this effect also reduced overexpression of *II-15* (Sun and Liu, 2015). I also confirmed associations with butter intake and pentadecanoate (15:0), 10-undecenoate (11:1n1) and 15-methylpalmitate from previous metabolomics studies (Guertin et al., 2014b, Zheng et al., 2014).

Low fat milk intake was strongly associated with a recently identified metabolite trimethyl-N-aminovalerate (X-21365). I replicated this association between general milk intake and trimethyl-N-animovalerate in the KORA population. The origin of this novel association is complicated to decipher as literature is limited on trimethyl-N-aminovalerate. Trimethyl-N-aminovalerate is potentially a methylated product of 5-aminovalerate, a product of lysine or proline degradation by gut bacteria *Escherichia coli* (Li et al., 2016) and *Pseudomonas putida* (Revelles et al., 2005). Moreover, both molecules are structurally similar to carnitine. Further research will be needed to unravel the mechanism of this association.

Grain-rich product intake

I found higher reported intakes of high fibre breakfast cereals to be associated uniquely with increased levels of pyridoxate, a vitamin B6 metabolite. Pyridoxate has many essential roles including being a coenzyme in the synthesis of amino acids, sphingolipids, neurotransmitters (serotonin, norepinephrine) and aminolevulinic acid. Higher pyridoxate in bloods has been associated with increased reported consumption of vitamins/supplements and other fruits (including plums, apricots, peaches, prunes, raisins, grapes and pineapple), as well as increased scores on the Healthy Eating Index (Guertin et al., 2014b). As it is common practice for breakfast cereals to be fortified with B vitamins this may have contributed to higher levels. I also found increased porridge intake to be associated with higher levels of 2-aminophenol sulphate. Previously, higher urinary output of 2-aminophenol sulphate was found in subjects fed whole grain rye bread versus refined wheat bread in a cross-over intervention study (Bondia-Pons et al., 2013). 2-aminophenol sulfate is a benzoxazinoid metabolite. Whole grains contain higher levels of benzoxazinoids which are readily absorbed from these products (Jensen et al., 2015). Higher reported intakes of refined grain products were associated with increased octenoylcarnitine. Octenoylcarnitine is an acylcarnitine formed from mitochondrial betaoxidation. A previous metabolomics study undertaken on 33 coeliac disease patients showed those following a long-term gluten free diet had reduced octenoylcarnitine (Bene et al., 2005).

Fruit consumption

Higher reported intakes of apples and pears were associated uniquely with a sugar alcohol, threitol, and two amino acids formed from gut bacterial metabolism of phenolic compounds: indolepropionate (also associated with bananas) (Smith and Macfarlane, 1996, Karbownik et al.,

2001) and 3-phenylpropionate (Smith and Macfarlane, 1996). Apple proanthocyanidins have shown to be directly catabolized to 3-phenylpropionate when incubated with human gut bacteria (Ou et al., 2014). A previous study showed higher reported intakes of eggs and red meat in a US population to associated with lower indolepropionate in blood (Guertin et al., 2014b). Their finding may suggest these subjects had lower intakes of fruit-derived proanthocyanidins or higher consumption of animal proteins, which have been shown to rapidly alter the gut microbiome composition (David et al., 2014), and hence may indirectly influence bacterial metabolism of proanthocyanidins through altering the intestinal environment. Previously, increased indolepropionate has been associated with insulin sensitivity (Gall et al., 2010), and lower levels associated with reduced muscle mass in elderly subjects (Lustgarten et al., 2014). For 3-phenylpropionate (ACSM5, rs11647589) and indolepropionate (ACSM2A, rs1394678) SNPs within medium-chain acyl-CoA synthetase (MACS) genes were associated with levels. The ligation of medium-chain fatty acids with CoA is catalysed by MACS genes, though MACS also conjugate glycine with benzoic acid derivatices (Kasuya et al., 1996). Similar to 3phenylpropionate and indolepropionate, benzoic acid is produced by gut microbial degradation of apple and cranberry phenolic compounds (Ou et al., 2014), the genotypic association with 3phenylpropionate and indolepropionate may be due to their correlation to products of this process. An association with metabolic syndrome phenotypes and a variant in the ACSM2 gene have been found previously; though this this association may be due mainly to the role of ACSM2 in lipid metabolism (Lindner et al., 2006). I confirmed a previous association between reported intakes of fruit juice with stachydrine (also proline betaine) (Guertin et al., 2014b, Zheng et al., 2014), a component found in citrus fruits.

Vegetable consumption

Reported mushroom intake was associated with ergothioneine, a novel finding. Ergothioneine is a thiol compound found in higher quantities in specialty mushrooms such as oyster and king bolete (Ey et al., 2007). Ergothioneine has also shown to prevent lipid peroxidation *in vivo* (Deiana et al., 2004) and recent data suggest that ergothioneine may prevent vascular dysfunction (Li et al., 2014). I found higher reported intakes of green/leafy vegetables and avocado to associated with higher levels of metabolites that are strongly associated with seafood consumption including CMPF (3-carboxy-4-methyl-5-propyl-2-furanpropanoate) and 1docosahexaenoylglycerophosphocholine, this issue may have arisen due to highly correlated

intakes, a problem encountered previously (Guertin et al., 2014a, Zheng et al., 2014, Ling et al., 2014).

Tea and coffee

I identified higher reported intakes of herbal tea were associated with lower levels of long-chain phosphatidylcholine acyl-alkyls derived from hepatic lipid metabolism (Wittenbecher et al., 2015). A dietary pattern characterised by a high intake of red meat and fish and lower intakes of whole grain bread and tea was previously shown to correlate with reduced levels of phosphatidylcholine diacyl (including phosphatidylcholine diacyl C42:4) in the EPIC Potsdam cohort (Floegel et al., 2013b). Also in the EPIC Potsdam cohort, levels of these phosphatidylcholines have associated with diabetes risk (Floegel et al., 2013a), though little is known currently about the origin of these associations.

I also confirmed associations between increased reported coffee intake and higher levels of metabolites derived from caffeine and coffee (Guertin et al., 2014a, Zheng et al., 2014). Interestingly, I also found a number of metabolites derived from coffee were associated inversely with reported black tea intake. This pattern is suggestive that individuals who reported having a higher tea intake had habitually lower coffee consumption. I identified higher levels of O-methyl catechol sulfate to be associated with higher reported coffee intake, which was a novel finding. One of the strongest metabolites associated to coffee intake was 1-methylxanthine. 1-methylxanthine is derived from caffeine metabolism and was previously associated to a variant in *NAT2* (rs4921914) which encodes a liver enzyme that acetylates caffeine metabolites (Butler et al., 1992). A SNP in *NAT2* has found to modulate an association with black tea consumption and risk of SLE (Kiyohara et al., 2014).

Sweet & savoury discretionary food intakes

Multiple associations were identified between foods where reduced consumption is promoted ('discretionary'; such as sweet baked products, sweets and jams, crisps, fried potatoes and fish and savoury pies) that seem to not be biologically sound and are also novel. This could suggest that these associations are showing habitual reduced intakes of other foods (i.e. vegetables, fruit, fish and wine) with increased reported intakes of discretionary foods. For instance, increased reported consumption of sweet baked products (including cookies, cakes, pies and pastries) were associated with lower levels of essential fatty acids derived from fish, (DHA and

EPA) and scyllo-inositol (strongly correlated with wine intake). Also, I found higher intakes of fried fish and savoury pies to be associated to lower levels of 3-phenylpropionate derived from gut microbial catabolism of fruit proanthocyanidins (Ou et al., 2014). Reporting of these foods may have been influenced by self-reporting biases (Westerterp and Goris, 2002), a limitation of this dataset.

Other notable associations

I identified a novel association with increased soymilk consumption and higher levels of 4ethylphenylsulfate, despite intakes being low in the sample; a previous study of US subjects found higher 4-ethylphenylsulfate to be correlated with increased tofu consumption (Guertin et al., 2014b). 4-ethylphenylsulfate is formed under gut bacteria metabolism. Elevated levels of 4ethylphenylsulfate have been found to correlate with anxiety-like traits in rats. Increased levels of 4-ethylphenylsulfate in blood are thought to be due to higher gut permeability (Hsiao et al., 2013). This association may be derived from the high saponin content of soybeans, which have shown to increase intestinal cell permeability in vitro (Johnson et al., 1986) though also in Atlantic salmon (Knudsen et al., 2008).

I found a novel association between higher reported chocolate intake and increased 7methylxanthine, a methylated purine, and I also confirmed an association with theobromine, a bitter alkaloid and recognised marker of cocoa intake (Rodriguez et al., 2015). 7-methylxanthine is derived from methylxanthine metabolism, methylxanthines include caffeine, theophylline and theobromine (Suzuki and Takahashi, 1975).

Notable unknown metabolite associations

The chemical identity of 31 metabolites associated to food intakes are currently unknown, though these may become important food intake biomarkers in the future. In particular I found an association between increased reported fried food intake (fried fish and savoury snacks [including potato crisps]) and higher levels of the metabolite X-11372. I also identified another metabolite associated positively with red and processed meat consumption, metabolite X-11381. Metabolite X-11381 was previously associated with a SNP in a gene encoding a carnitine efflux transporter *SLC16A9* (rs12356193) (Suhre et al., 2011). Variants in *SLC16A9* have previously been associated with serum uric acid levels (Lee and Song, 2012) and risk of gout in renal overload (Nakayama et al., 2013). Another unknown metabolite, X-09789, was

associated positively with porridge intake. A variant in the *SLC51A* gene (also known as *OST-alpha*; rs7642243) was found to be associated with X-09789. Interestingly, *SLC51A* encodes a component of the Ost-alpha/Ost-beta complex which functions in bile acid transport from intestinal enterocytes into portal blood (Ballatori et al., 2008). Oats are known to contain beta-glucan, a soluble fibre found to reduce blood cholesterol levels (Charlton et al., 2012) by sequestering intestinal bile acids and effectively reducing the reabsorption of bile acids (Wolever et al., 2010). I also found the unknown metabolite X-11315 to be associated with reported consumption 13 foods (top association: apple and pears; negative associations with discretionary foods). Metabolite X-11315 was previously associated to a SNP in *SLC6A20* (rs4327428), which encodes a proline transporter, this suggests metabolite X-11315 is structurally like proline. Previous studies have identified polymorphisms in *SLC6A20* to be associated with risk of Type 2 diabetes in white-European and Chinese populations (Ling et al., 2014).

4.4.2 Dietary patterns discussion

Examining dietary patterns in conjunction with metabolomics, I identified a number of potential metabolite biomarkers of dietary intakes in the twin population and validated the results using the co-twin control method. The repetitive occurrence of a number of metabolites within opposing dietary patterns in this study (**Table 4-5**) further confirms the utility of metabolomics for the identification and exploration of diet and disease relationships. I also identified a number of metabolites unique to each diet pattern (**Table 4-6**). Many of the associations with dietary patterns were attributable to their association with food intakes, therefore I will not discuss these associations in detail. However, there were 11 known metabolites and 10 unknown metabolites independently. The Traditional English and Dieting patterns did not appear to have a substantial metabolomic footprint and did not capture unique associations with metabolites, this may be due to underreporting.

Fruit and vegetable pattern

Twins scoring highly on the F&V pattern presented with an increase in metabolites relating to antioxidant potential, including hippurate which accumulates following increased consumption of

foods containing high phenolic compounds such as green and black teas (Van Dorsten et al., 2006); threonate, a product of vitamin C metabolism; and the γ -glutamyl peptide, γ -glutamylvaline and proline which both suggest reduced glutathione and collagen breakdown on this diet. Higher scores on the F&V diet pattern were also associated with increased phosphatidylcholine diacyl 38:6 and N-acetylornithine. N-acetylorthinine is an endogenously synthesized amino acid involved in urea metabolism. A genetic variant in N-acetyltransferase 8 gene (*NAT8*) was found to contribute to N-acetylornithine levels which may play role in kidney function and CKD (Yu et al., 2014). Despite the MDS and F&V patterns characterised by similar foods the data-driven F&V score captured unique metabolite associations, suggesting that measuring a Mediterranean diet in a non-Mediterranean country such as the UK may be problematic, an issue which has been thoroughly discussed previously (Hoffman and Gerber, 2013).

Low meat dietary pattern

Higher scores of the low meat dietary pattern were associated with lower levels of phosphatidylcholine acyl-alkyl 38:6 and higher levels of betaine and C-glycosyltryptophan. Betaine is an amino acid derived from choline. Choline is a nutrient that has an important role in methylation processes and reduction of homocysteine levels and subsequent risk of coronary heart disease (Zeisel and da Costa, 2009), the origin of this association requires further investigation. The low meat diet pattern was positively associated with C-glycosyltryptophan. Circulating levels of C-glycosyltryptophan have been associated with aging and age-related diseases in the TwinsUK population (Menni et al., 2013b), negatively associated with muscle mass in less mobile elderly subjects (Lustgarten et al., 2014), and a CpG site in the promoter of the WDR85 gene, which is required for protein synthesis (Menni et al., 2013b). This could suggest that individuals consuming a dietary pattern low in protein but high in starches and carbohydrates tend to be older or this type of dietary pattern may contribute to physical aging and reduced muscle mass.

High alcohol dietary pattern

A unique association was identified between the high alcohol diet and 4-vinylphenol sulphate, a product of benzoate metabolism and compound resulting from nut roasting (Walradt, 1971), and airborne styrene exposure and cigarette smoking (Manini et al., 2003), that has been previously
associated with nut consumption (Guertin et al., 2014b) and several CpG loci (Petersen et al., 2014). The metabolic implications for this epigenetic association are unknown at this time, though it may suggest that individuals scoring highly on this pattern are more prone to cigarette smoking. Increased scores on the alcohol pattern were also associated with higher levels of gamma-tocopherol, Gamma-tocopherol is one form of vitamin E, a major dietary antioxidant. The relationship between alcohol intake and gamma-tocopherol is unclear, a previous study found higher intakes of vitamins and supplements, and scores on the US Healthy Eating Index were associated with lower serum γ -tocopherol (Guertin et al., 2014b).

Table 4-5. List of	metabolites	associated to	o multiple	dietary	patterns

Super-pathway	Sub-pathway	Metabolite	MDS	F&V	High alcohol	Traditional English	Low meat
Amino acid	Creatine metabolism	Creatine				1	\downarrow
	Phenylalanine & tyrosine metabolism	3-Phenylpropionate (hydrocinnamate)		1		Ļ	
	Tryptophan metabolism	Indolepropionate	1	↑		\downarrow	
	Valine, leucine and isoleucine metabolism	α-Hydroxyisovalerate			1		\downarrow
Carbohydrate	Glycolysis, gluconeogenesis, pyruvate metabolism	1,5-Anhydroglucitol (1,5-AG)		\downarrow			ſ
		Glycerate	1	1			
Lipid	Essential fatty acid	Docosahexaenoate (DHA; 22:6n3)	ſ	ſ			\downarrow
		Eicosapentaenoate (EPA; 20:5n3)	1	<u>↑</u>			\downarrow
	Fatty acid, dicarboxylate	3-Carboxy-4-methyl-5-propyl-2-furanpropanoate (CMPF)	ſ	1			\downarrow
	Inositol metabolism	Scyllo-inositol	1	<u>↑</u>	<u>↑</u>		\downarrow
	Lysolipid	1-Docosahexaenoylglycerophosphocholine	1	↑			
	Medium chain fatty acid	10-Undecenoate (11:1n1)			\uparrow		\downarrow
Xenobiotics	Benzoate metabolism	Catechol sulfate		1		\downarrow	

Dietary phenotype	Super-pathway	Sub-pathway	Metabolite	Direction of association
Fruit & vegetable	Amino acid	Lysine metabolism	Pipecolate	↑
J. J	Carbohydrate	Nucleotide sugars, pentose metabolism	Threitol	1
	Cofactors and vitamins	Ascorbate and aldarate metabolism	Threonate	1
	Peptide	gamma-glutamyl	γ-Glutamylvaline	Ļ
	Xenobiotics	Benzoate metabolism	Hippurate	1
High alcohol	Amino acid	Butanoate metabolism	2-Aminobutyrate	1
	Lipid	Inositol metabolism	Myo-inositol	↑
		Medium chain fatty acid	Caprate (10:0)	1
		Medium chain fatty acid	10-Undecenoate (11:1n1)	↑
	Xenobiotics	Food component/Plant	Piperine	↑
Traditional English	Amino acid	Urea cycle; arginine-, proline-, metabolism	Stachydrine	Ļ
		Urea cycle; arginine-, proline-, metabolism	Trans-4-hydroxyproline (Hydroxyproline)	1
Dieting	Xenobiotics	Benzoate metabolism	4-Ethylphenylsulfate	1
-		Food component/Plant	Quinate	Ļ
Low meat	Amino acid	Glutamate metabolism	Pyroglutamine	1
		Glycine, serine and threonine metabolism	Betaine	1
		Tryptophan metabolism	C-Glycosyltryptophan	1
		Cysteine, methionine, SAM, taurine metabolism	2-Hydroxybutyrate (AHB)	\downarrow

Table 4-6. List of metabolites associated to specific dietary phenotype

Strengths and Limitations

Although I replicated many associations from previous diet-metabolomics studies, there were multiple limitations to my study. The population I used was only female and as such my findings may not apply to men. Replicating my top associations in an independent population would have strengthened my findings, although I used the twin model for validation, allowing me to use controls matched for age, sex and the baseline genetic sequence. I did however, confirm associations from similar studies (Guertin et al., 2014b, Zheng et al., 2014) which suggests the data used in the study were of decent quality. Due to the cross-sectional nature of this study design, the study does not allow me to indicate cause and effect. As I used only FFQ data as my source for dietary information, the accuracy of the data can be called into question due to the consequences of self-reporting (Westerterp and Goris, 2002). Despite this issue, many of the associations I identified are biologically plausible. Moreover, food frequency data only provides categorical information and therefore the precise effect cannot be quantified, though this is an area where future dietary intervention studies can aid findings. I do recognize I may have experienced a small number of associations due to type 1 errors potentially as a result of correlated intake reporting, this issue may have been encountered for foods which lower intakes are generally discouraged, including sweet and savoury discretionary foods (sweets and jams, sweet baked products, fried fish and savoury pies). Using very stringent multiple-testing cut-offs I am hopeful many of these spurious associations were minimized, although this likely caused false negatives. Moreover, I did not have longitudinal data on metabolite levels, which in future studies could provide useful information on metabolite stability over time and may improve the strength of associations with food intakes. The true blood concentration of the metabolites is not measured by the Metabolon platform, therefore accurate quantification could not be determined for these metabolites. Furthermore, the current study only provides a snapshot of associations in one type of tissue, at only one time point and therefore provides limited insight into the origin and directionality of associations and virtually no insight into mechanisms. Having genotypic profiling completed on a large sample of TwinsUK with results also replicated in KORA, this allowed me to supplement my discussion, though future studies would need to untangle these potential genotype-metabolite-disease relationships and whether they are modulated by dietary factors. Strengths of this chapter include the large sample size and number of metabolites that were assessed and I was also able to use MZ discordant twins who are matched for age, sex and the baseline genetic sequence.

Conclusions

By using one of the largest and comprehensive datasets of its kind, I identified 178 self-reported food intake associations (72 novel) with blood metabolites. I also identified 108 associations with 6 dietary patterns, confirming dietary patterns are associated with a unique metabolomic signature not captured by assessing food intakes independently. Future studies should aim to undertake dietary interventions trials to confirm our findings, adequately determine mechanisms for associations and quantify the effect of food intake on metabolite levels. The findings of my study can be viewed online using the DietMetab search tool

(http://www.twinsuk.ac.uk/dietmetab-data/).

Chapter 5 Creating metabolite scores based on reported food group intake and applying these to explore the impact of diet on the metabolic syndrome

In this chapter I analyse 20 food groups for their association with metabolites from the nontargeted (Metabolon) platform, I then use three methods to combine multiple metabolites associated with food group intakes and evaluate them in an independent testing dataset. Therefore the overall aim of this chapter is to assess combining metabolites as a means to strengthen the utility of food intake biomarkers. Finally, I use the top performing metabolite scores to evaluate their correlation with an example of a common disorder - metabolic syndrome risk.

5.1 Introduction

In the previous chapter I showed that reported food intakes are associated with a unique blood metabolomic fingerprint. In order to best categorise habitual food intake to improve disease risk prediction, the appropriate handling of identified biomarkers must first be evaluated.

Used alone single biomarkers may be less precise as they can be influenced by a multitude of factors and non-specific to a particular disease state. For example, c-reactive protein is synthesised by the liver under inflammatory states and is therefore used as a general marker for inflammatory processes occurring in the body and not specific to one condition. Indeed it is one marker of the metabolic syndrome (Olza et al., 2015) and CVD risk (Berezin et al., 2015). Moreover, complex chronic diseases are characterised by clusters of symptoms that cannot be quantified by measuring a single biomarker. Biomarker risk scores have been evaluated and shown to improve accuracy over traditional methods for a multitude of conditions, such as CVD (Berezin et al., 2015, Richter et al., 2013, Hughes et al., 2012) and metabolic syndrome (Olza et al., 2015).

Similarly, combining dietary information to create scores is currently performed, such as for creating dietary patterns *a priori* to measure adherence to dietary patterns associated with particular health outcomes or measuring diet quality, including the MDS (Trichopoulou et al., 2003) and the Healthy Eating Index (Guenther et al., 2013). Scoring is assigned using cut-offs

points previously associated to health outcomes or diet quality. Diet quality is indicated by scoring reported food intakes by how closely they adhere to established dietary guidelines and also the variety of healthy food choices within core food groups. Dietary adherence scores have been tested for their correlation with traditional biomarkers of nutrient intake (e.g. n-3 fatty acids and carotenoids) (Golley et al., 2015). A study from EPIC-Norfolk created a panel of traditional biomarkers of fruit and vegetable intakes by summing plasma vitamin C, beta-carotene and lutein and found the panel to be strongly inversely associated with incident type II diabetes (Cooper et al., 2015). Moreover, a multi-food intake biomarker approach applying summation and PCA methods was used to evaluate adherence to a healthy Nordic diet dietary intervention in patients with MetS (Marklund et al., 2014), both methods performed similarly. With emerging nutritional metabolomics studies measuring multiple biomarkers simultaneously these traditional biomarkers may soon be replaced by more precise food intake scores.

In one nutritional metabolomics study biomarkers of alcohol consumption in African Americans were identified and combined to create an alcohol risk score to evaluate the effects of alcohol consumption on levels of white blood cells and incident CVD (Zheng et al., 2014) . Specifically, the authors created three composite scores by summing quartile-ranked metabolites belonging to the same alcohol-related metabolic sub-pathways, including 2-hydroxybutyrate and related BCAAs, lysophosphatidylcholines, and γ-glutamyl dipeptides. However, combining this information may have been more useful as independently these pathways are affected by different health conditions. In another study, a metabolomics approach was used to identify a panel of metabolites related to sugar-sweetened beverage (SSB) consumption, where a panel of 4 metabolites (formate, citrulline, taurine, and isocitrate) were found to be discriminate consumers and non-consumers (Gibbons et al., 2015).

Combined biomarker scores representing long-term intakes of frequently consumed foods may greatly improve our understanding of the health impacts of diet. Therefore, my aims for this chapter were to:

- i) Identify potential biomarkers of intakes of 20 commonly consumed food groups.
- ii) Form food metabolite scores using multiple methods and test their utility.
- iii) Use the scores to predict the risk of developing the metabolic syndrome.

5.2 Materials and methods

Figure 5-1 shows the pipeline of the chapter.



5.2.1 Food groups

I created and analysed twenty food groups of similar taste and nutrient content (shown in Table

5-1) derived from FFQ against blood metabolites.

Table 5-1. List of food items included in food groups

Food group	FFQ items
Vegetables	
	Broccoli, spring green, kale
	Brussel sprouts
	Cabbage
	Cauliflower
	Coleslaw
	Avocado
	Beetroot
	Marrow, courgettes
	Mushrooms
	Parsnips, turnips, swedes
	Sweetcorn
	Sweet peppers
	Watercress
	Carrots
	Tomatoes
	Garlic (clove)

Food group	
rood group	
	Leeks
	Onions
	Green salad, lettuce, cucumper, celery
	Spinach
	Watercress
	Vegetable soups (bowl)
	Boiled, mashed, instant or one jacket potato
Fruit	
	Strawberries, raspberries, other berries, kiwi fruit (one fruit or handful)
	Smootnies (cup)
	Pure fruit Juice (100%) e.g. orange, apple Juice (cup)
	Oranges, satsumas, mandarins (1 fruit)
	Apples (1 fruit)
	Bananas (1truit)
	Dried fruit, e.g. raisins, prunes (heaped tablespoon)
	Grapes (handful)
	ivieron (1 sliče) Deselves sluves survisets (4 furit)
	Peaches, plums, apricots (1 truit)
	Pears (1 muit)
M/halss -	linnea truit (nanatul)
whole grains	Ligh Fibre corrects on a Dranflakes. All Dran, Envit and Fibre
	nigh Fibre Cereais e.g. Brannakes, All Bran, Fruit and Fibre Mueeli
	Muesii Derriden Derdeharste este
	Porriage, keadybreak, oats
	Brown rice
	Wholemeal agent
Defined areing	Crisporead, e.g. Ryvita
Renned grains	Proakfast coroal o g. Coroflakos. Dico Krisnics
	Sugar topped coreals on a Fraction
	Naan nonnadoms flour tortillas
	Rown broad/rolls
	White broad /rolls
	White or groop pasta, o.g. spaghotti, macaroni, poodlos
	White rice
Nuts and logu	mac
Nuts and legul	Boonsprouts
	Dulsas a glantils haans naas
	Green heans broad heans runner heans
	Dese
	r cas Raked heans
	Salted nuts e.g. neanuts, cashews (handful)
	Linsalted nuts e.g. peanars, cashews (nanarar)
	Seeds e g Sunflower numnkin (tablesnoon)
	Peanut hutter (teasnoon)
	Meat substitutes e gi tofu sovameat texturad vegetable protein vegeburger
Seafood	אוכער שעשטורעונט כ.ב. נטוע, שטימוויכער, ובאנערכע עכצבומטוב אוטוכווו, עכצבטעוצבו
Jearoou	Oily fish, fresh or canned, e.g. tuna, mackerel, kinners, salmon, sardines, herring
	Fish roe, taramasalata
	Shellfish, e.g. crab, prawns, mussels
	Other white fish, fresh or frozen, e.g. cod plaice sole haddock halibut
White meat	
e meat	Chicken or other poultry e.g. turkey
Red meat and	eggs
	Beef: roast, steak, mince, stew or casserole
	Lamb: roast, chops or stew
	Pork: roast, chops or stew
	Beefburgers
	Bacon or gammon
	Corned Beef, Spam, luncheon meats
	Ham cured meats & chorizo

Table 5-1. List of food items included in food groups

Table 5-1. List	of food items	included	in food	groups
-----------------	---------------	----------	---------	--------

Food group FFQ items	
Liver, liver pate, liver sausage	
Sausages	
Eggs as boiled, fried, scrambled, etc. (one)	
Fermented dairy	
Low fat cheese e.g. reduced fat cheddar (matchbox size)	
Cheese, e.g. cheddar, brie, edam (matchbox size)	
Cottage cheese, low fat soft cheese (2 tablespoons)	
Full fat or Greek yoghurt (small pot)	
Low fat yoghurt, fromage frais (small pot)	
Fried foods	
Fish fingers, fish cakes & breaded fish	
Fried fish in batter, as in fish and chips	
Chips, retail, fried in vegetable oil	
Potato salad	
Old potatoes, roast in blended oil	
Savoury pies, e.g. meat pie, pork pie, pasties, steak & kidn	ey pie, sausage rolls
Cream crackers, savoury biscuits	
Crisps or other packet snacks, e.g. Wotsits (one packet)	
Pizza (one slice)	
Quiche (slice)	
Sweets and sweet baked products	
Reduced fat biscuits e.g. Go Ahead, Highlights (one small p	backet or one small bar/biscuit)
Sweet biscuits, chocolate, e.g. digestive (one)	
Sweet biscuits, plain, e.g. Nice, ginger (one)	
Buns, pastries e.g. scones, flapjacks, croissants, doughnuts	s, home baked
Cakes e.g. fruit, sponge, home baked	
Cakes e.g. fruit, sponge, ready made	
Fruit pies, tarts, crumbles, home baked	
Fruit pies, tarts, crumbles, ready made	
Milk puddings e.g. rice, custard, trifle	
Sponge puddings, nome baked	
Sponge puddings, ready made	
Dairy desserts (small pot) e.g. chocolate mousse, cream ca	aramels
Ice cream, choc ices	
Jam, marmalade, honey (teaspoon)	
Sugar added to tea, coffee, cereal (teaspoon)	
Sweets, torrees, mints (smail packet)	
Chocolate	
Dark chocolates, single or squares (one)	
while of milk chocolates, single of squares (one)	
Low fat not chocolate (cup)	
Cocoa, not chocolate (cup) Chocolate spack bars o a Mars. Crunchia (ano)	
Butter and cream	
Butter and cream Reduced fat hutter (teaspoon)	
Butter (teaspoon)	
Double or clotted cream (tablespoon)	
Single or sour cream (tablespoon)	
Spreads and dressings	
Low fat spread, e.g. Outline, Gold (teaspoon)	
Very low fat spread (teaspoon) e.g. Diet Flora	
Cholesterol lowering fat spreads e.g. Benecol (teaspoon)	
Olive oil spread (teaspoon)	
Block margarine e.g. Stork Krona (teasnoon)	
Other soft margarine, dairy spreads, e.g. Blue Band, Clove	r (teaspoon)
Polyunsaturated margarine, e.g. Flora, sunflower (teasnor	
French dressing (tablesnoon)	
Full fat salad cream, mayonnaico (tablosnoon)	
Other salad dressing (tablespoon)	
Low calorie low fat salad cream (tablespoon)	
Milk	
Channel Islands milk	
Full cream milk	

Food group	FFQ items
<u> </u>	Dried milk
	Semi-skimmed milk
	Skimmed milk
Soya and othe	r milk
	Goats' milk
	Rice milk
	Soya milk
Soda	
	Fizzy soft drinks, e.g. Coca Cola, lemonade (cup)
	Low calorie or diet fizzy soft drinks (cup)
Теа	
	Tea (cup)
	Green tea (cup)
Coffee	
	Coffee, instant or ground (cup)
	Coffee, decaffeinated (cup)
Alcohol	
	Beer, lager or cider (half pint)
	Port, sherry, vermouth, liqueurs (pub measure)
	Spirits, e.g. gin, brandy, whisky, vodka (pub measure)
	Red wine (small glass)
	White wine (small glass)

Table 5-1. List of food items included in food arouns

5.2.2 **Metabolomics dataset**

Fasted blood metabolites derived from the Metabolon dataset were used (Section 3.1.6.1). I included in the analysis 329 metabolites that were present in both serum and plasma and had no missing values in 75% of the sample.

5.2.3 Training and validation datasets

Female twins with Metabolon data available within and including 5 years of FFQ completion were included in the total sample (n=3559). A subset of 1156 twins had food preference data available, I therefore randomly assigned 1779 twins (keeping co-twins together) of this subset (50% of the whole sample) to the validation group, while the remaining 50% of the sample (n=1780) were assigned to the training group. Table 5-2 shows the characteristics of the training and validation datasets.

Table 5-2. Characteristics of the training and validation groups

		Training	group		Valio	dation group
	D	iscovery ¹	M	Z discordant		
Phenotype	n	Mean (SD)	n	Mean (SD)	n	Mean (SD)
Age (years)	1780	56.9 (14.9)			1779	54.4 (11.6)
BMI (kg/m2)	1766	26.1 (4.8)			1771	26.1 (4.9)
Food groups (servings/week) ²						
Vegetables	1652	32.1 (16.2)	128	40.4 (24.1)	1779	34.2 (15.5)
Fruit	1588	20.3 (12.7)	192	26.2 (16.9)	1779	21.8 (12.5)
Whole grains	1582	9.7 (8.6)	198	14.4 (11.2)	1779	10.3 (8.2)

	Training group Va					lation group
	D	iscovery ¹	MZ	discordant		
Phenotype	n	Mean (SD)	n	Mean (SD)	n	Mean (SD)
Refined grains	1588	10 (8.3)	192	13.9 (11.2)	1779	9.6 (8.2)
Nuts and legumes	1598	6.5 (4.3)	182	9.9 (6.1)	1779	7.5 (5.3)
Seafood	1590	2.1 (1.8)	190	3.5 (2.7)	1779	2.4 (1.9)
White meat	1492	1.8 (1.3)	288	2.2 (1.4)	1779	2.0 (1.3)
Red meat	1626	6.5 (3.7)	154	8.5 (5.1)	1779	6.7 (3.8)
Fermented dairy	1618	5.5 (4.3)	162	8.2 (7.2)	1779	6.2 (4.8)
Fried foods	1604	4.9 (3.6)	176	7.3 (5.9)	1779	4.5 (3.3)
Sweets and sweet baked products	1618	17.4 (14.5)	162	27.0 (22.6)	1779	15.6 (14)
Chocolate	1662	3.3 (5.0)	118	7.8 (9.3)	1779	4.0 (6.0)
Butter and cream	1616	3.3 (5.7)	164	8.6 (8.8)	1779	4.1 (6.4)
Spreads and dressings	1610	8.8 (8.5)	170	14.9 (13.2)	1779	8.6 (9.1)
Milk	1592	3.7 (2.2)	188	4.4 (3.1)	1779	3.4 (2.2)
Soya and other milk	1752	0.1 (0.6)	28	2.0 (2.1)	1779	0.2 (0.8)
Soda	1672	2.3 (5.2)	108	7.8 (8.3)	1779	2.3 (5.5)
Теа	1584	20.2 (13.8)	196	19.7 (13.8)	1779	19.0 (13.8)
Coffee	1574	8.9 (10.8)	206	15.9 (12.6)	1779	8.9 (10.7)
Alcohol	1682	4.9 (7.1)	98	12.0 (11.7)	1779	6.3 (7.8)

Table 5-2. Characteristics of the training and validation groups

¹Excluding twins discordant for food group intake.

²Variables are residual adjusted for energy intake.

5.2.4 Classification of the metabolic syndrome

During clinical visits undertaken on the twins estimates of waist circumference were determined by analysing DXA scans, fasted blood samples were taken and analysed to determine levels of total triglycerides (TG), high density lipoprotein (HDL)-cholesterol and glucose, and blood pressure (BP) was determined, consult **Section 3.5.1** for details.

I determined MetS status using the criteria outlined by the International Diabetes Federation and the American Heart Association/National Heart, Lung, and Blood Institute (Alberti et al., 2009). Twins who had at least three of any of the following factors were considered to have the metabolic syndrome:

- Increased waist circumference (WC): men ≥94 cm and women ≥80 cm, or BMI > 30 kg/m².
- 2. Increased triglycerides (TG): >1.7 mmol/L, or treatment for abnormality
- Lower HDL-cholesterol: men <1.03 mmol/L in men and women <1.29 mmol/L, or treatment for abnormality
- 4. High blood pressure: ≥130/85 mm Hg, or treatment for hypertension
- Raised fasting plasma glucose (FBG): FBG ≥ 5.6 mmol/L. Most individuals with type 2 diabetes will have metabolic syndrome based on these criteria

I included 414 twins in the analysis who had metabolomics profiling and diet information and did not have MetS at this time. These twins also attended a clinical visit 5 years or more after metabolomics profiling where MetS status was confirmed.

5.2.5 Statistical analysis

Statistical analysis was carried out using Stata version 12.

5.2.5.1 Discovery analysis in the training dataset

For each metabolite, random intercept linear regression analysis was undertaken in the first sample (discovery sample) excluding MZ twin pairs discordant (MZ twins with measures one SD apart in food group intake) for each food group. Age, metabolite batch, BMI and family relatedness were included as covariates:

$$Y_{i} = \beta_{0} + \beta_{i} X_{ii} + \gamma_{i} age_{ii} + \delta_{i} BMI_{ii} + \zeta_{i} + \varepsilon_{ii}$$

where Y_i is the metabolite and X_{ij} the food group intake of twin *j* from pair *i*, ζ_j , is the family-specific error component that captures the unobserved heterogeneity or family characteristics.

I adjusted for multiple testing using Bonferroni correction thus giving a significant threshold of 7.60x10⁻⁶ (0.05/(20 food groups x 329 eligible metabolites)). For each significant metabolitefood group association from the discovery sample, I repeated the same linear regression analysis on the MZ discordant twin pair sample. The MZ discordant twin pairs were used to replicate the significant findings from the discovery group, associations in the same direction as the discovery group were considered replicated. Finally, we combined the results of both analyses using an inverse variance fixed effects meta-analysis that are the reported results. The beta coefficients (β) presented in the results of each linear regression analysis represent the amount of a food group consumed in servings per week that corresponds to a 1 SD change in the metabolite level.

5.2.5.2 Data reduction

Following identification of significant metabolites for each food I reduced the number of associations.

Firstly, a number of metabolites were associated with multiple food groups, which may have been a result of correlations in intake reporting. I therefore conducted a backward stepwise linear regression using the 5% level of significance as the cut-off threshold for each metabolite (adjusted for covariates) using all food groups associated as predictors of the

metabolite accordingly. I considered associations to be significant if they passed the Bonferroni cut-off for multiple testing used for the discovery analysis (P<7.60x10⁻⁶).

Secondly, a number of the food groups were associated with multiple potentially correlated metabolites. Although the hypothesis is that metabolite levels change with food intake, to determine which metabolites are correlated independently of one another with food intake I undertook a backwards stepwise linear regression (cut-off threshold: P<0.05), with food group intake as the dependent variable. I performed this on all the metabolite residuals identified following the discovery and first data reduction steps, to identify a potential panel of independent metabolites to use as food intake markers.

5.2.5.3 Food-metabolite group generation

Significant food group-metabolite associations were then converged into scores based on three different methods: 1) Similar to the method used by Zheng and colleagues (Zheng et al., 2014), I quartile ranked blood levels of significant metabolites and assigned the quartiles a score of 0 to 3 according to the direction of the association (i.e. positive association: Q1=0, Q2=1, Q3=2, Q4=3; negative association: Q1=3, Q2=2, Q3=1, Q4=0) and summed these values; 2) Like the method used by Marklund and colleagues (Marklund et al., 2014), I created a continuous summed score by summing the relative levels of each significant metabolite. Negatively-associated metabolites were multiplied by -1; 3) I created a weighted score by multiplying the relative levels by the beta-coefficients for each metabolite as a predictor of food intakes (standardized to have mean 0 and SD 1), adjusting for all other associated metabolites and summed the final scores. This method has been used by Dash and colleagues (Dash et al., 2013) to examine the indicators of oxidative balance and incident, sporadic colorectal adenomas.

5.2.5.4 ROC analysis by food intake

To evaluate the utility of the food metabolite scores as potential markers of food intakes I conducted a binary classification test in the validation group. I classified twins in lower tertile of each food group as a negative outcome (0), and the top tertile of food group intake I considered a positive outcome (1). The ability of the metabolite scores to correctly classify twins consuming high intakes (sensitivity; true positive rate) and correctly classify twins consuming low intakes (specificity; true negative rate) of the model was predicted and the receiver operating characteristic curve (ROC) generated by plotting the true positive rate against the false positive

rate at multiple threshold settings. I tested whether any scoring method had a consistent advantage over another method by testing the difference in the area under the ROC curve (AUC). The AUC may be interpreted as follows: AUC>0.9 has high accuracy, AUC>0.7–0.9 has moderate accuracy, AUC>0.5–0.7 has low accuracy, and AUC=0.5 is a chance result (Fischer et al., 2003).

Confirmation of Associations by Food Preference

I used an alternative form of getting diet information from subjects by asking them their food preferences. I adapted the food preference questionnaire developed in the US by Dr Valerie Duffy (Duffy et al., 2007) for use in the UK twin cohort (details in **Appendix D Document 1**). To do this, in collaboration with other scientists, items were changed to suit the UK diet and I transformed the questionnaire into an online format, this work has been published (Pallister et al., 2015). Modifications to the questionnaire involved changing the terms for some items (e.g. 'French fries' to 'chips', 'breakfast sausage' to 'sausage', and 'unflavoured oatmeal' to 'porridge') and some items were removed as they are infrequently consumed in the UK (e.g. 'bologna', 'canned noodle soup', 'chocolate milk', and 'blackened hot dog'). I compared their preferences for foods (ranked on a scale from -100 to +100) against those food groups as a way to further validate the metabolite scores. **Table 5-3** shows the foods which ratings were averaged to form the group. I ran a linear regression with the metabolite score as the response variable (adjusted for batch effects, BMI and age) and twin food group liking as the predictor variable adjusting for family relatedness. Those associations passing the cut-off for multiple testing (Bonferroni $P < 3.33 \times 10^{-3}$) were considered validated.

Table 5-3. List of food	preference items	by i	food	group	р
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Food group	Preference item
Vegetables	
	White potato
	Broccoli
	Fresh tomatoes
	Fresh coriander
	Garlic
	Sautéed mushrooms
	Asparagus
	Raw carrots
	Beetroot
	Spinach/ greens
	Aubergine
	Raw onion
Fruit	
	Grapefruit
	Banana
	Lemon

Table 5-5: Eist 6	rioou preference items by food group
Food group	Preference item
	Pear
	Melon
	Charries
	Chemies
	Strawberries
	Pineapple
	Orange juice
Whole grains	
U	Wholemeal bread
	Porridge
	Light fibro har
- C .	
Refined grains	
	Bagel/ rolls
	Pasta/ noodles
	Cornflakes
	White rice
Nuts and legumes	
Nuts and regames	Linealtad puts
	Faux meat products
	Lentils/ beans
Seafood	
	Tuna or salmon
	Prawns and shellfish
White meat	
	Baked chicken
Red meat	
Reumeat	Crimy hasan
	Crispy bacon
	Sausage
	Pork chops
	Ham
	Chargrilled meat
	Faas
	Boof stook
Fermented dairy pro	ducts
	Plain yoghurt
	Blue cheese
	Cheddar cheese
Fried and fast foods	
	Tortilla chips or crisps
	Chins
	Fried fish
	Pizza
Sweets and sweet ba	aked products
	lce cream
	Cake icing
	lam and jelly
	Riscuits cakes or nastries
	Chanceacha
0	
Chocolate	
	Dark chocolate
Butter	
	Butter or margarine
Spreads and dressing	25
	Butter or margarine
	Mayonnaise
	Salad dressing
	Extra virgin olive oil
Milk	
	Skimmed milk
	Whole milk
Soy milk	
	Sov milk

Table 5-3. List of food preference items by food group

Food group	Preference item
Soda	
	Carbonated drinks & sweet drinks
	Diet carbonated soft drinks
Теа	
	Hot tea
Coffee	
	Black coffee
	Coffee or tea with sugar
	Sweet coffee drinks & whipped cream
Alcohol	
	Beer
	Red wine
	White wine
	Vokda, gin or scotch

Table 5-3. List of food preference items by food group

5.2.5.5 Usage of food metabolite scores to predict the metabolic syndrome

To determine whether food metabolite scores were associated with risk of developing MetS, I used the food metabolite scores that performed the best (weighted scores if multiple metabolites and continuous variable for foods associated to single metabolites). I undertook a logistic regression in 414 twins using each food metabolite score (residual adjusted for age, BMI and batch at time of metabolomics) to predict future MetS status (0, no MetS [n=389]; 1, MetS [n=25]) adjusted for age, BMI and family relatedness. Statistical significance was defined as P<0.05. I performed the same logistic regression on reported food intake for metabolite scores significantly predictive of future MetS. I finally performed a logistic regression using each significant metabolite score as a predictor of each of the 5 clinical criteria for MetS.

5.3 Results

In the following paragraphs I will first present those metabolite associations used to create the food metabolite scores from the training dataset, followed by the validation results.

5.3.1 Overall associations with food group intakes

A summary of the results for the discovery and data reduction analysis can be found in **Table 5-4**. Following the meta-analysis of the discovery and discordant twin groups, 112 significant associations were identified in the training dataset (**Figure 5-2**; **Appendix D Table 1**). Significant associations were identified for all food groups except chocolate and refined grain intake. Thirty-nine metabolites were associated only with one food group (unique associations; **Table 5-5**), whereas 26 metabolites were associated with multiple foods (73 associations overall) potentially as a result of correlated intakes. Following a multivariate backward stepwise linear regression using each associated food group as a predictor of the latter 26 metabolites (**Appendix D Table 2**), 35 associations (including 24 metabolites) remained significant after Bonferroni correction (*P*<7.60x10⁻⁶).

Food group	Total metabolites associated in discovery analysis ⁽¹⁾	No. metabolites removed after 1 st data reduction ⁽²⁾	No. metabolites removed after 2 nd data reduction ⁽³⁾	Final no. metabolites	R ²
Vegetables	3	2	0	1	0.0094
Fruit	10	2	4	4	0.1195
Whole grains	7	4	0	3	0.0508
Nuts and legumes	2	1	0	1	0.0384
Seafood	18	3	8	7	0.1535
White meat	1	0	0	1	0.0291
Red and processed meat, and eggs	4	1	0	3	0.0403
Fermented dairy	1	1	0	0	-
Fried foods	5	4	0	1	0.0668
Sweets and sweet baked products	12	8	0	4	0.0588
Butter and cream	5	0	0	5	0.0754
Spreads and dressings	3	0	1	2	0.0331
Milk	5	2	0	3	0.0735
Soy and other milks	1	1	0	0	-
Soda	1	1	0	0	-
Теа	7	2	0	5	0.2113
Coffee	10	2	6	2	0.1963
Alcohol	17	4	5	8	0.2493

Table 5-4. Results summary of the discovery analysis, the 1^{st} and 2^{nd} data reductions, and the final number of metabolites contributing to each food-metabolite score with R^2

(1) Number of metabolites significantly associated to each food group in the discovery (Bonferroni; P<7.60x10⁻⁶) and the discordant MZ twin samples (P<0.05), and following fixed effects metaanalysis of results from both groups (Bonferroni; P<7.60x10⁻⁶).

(2) Metabolites with multiple associated foods were included in a backwards stepwise regression (cut-off threshold *P*<0.05), those associations not passing the significance threshold from the discovery analysis (*P*<7.60x10⁻⁶) were removed.

(3) Food groups associated with multiple metabolites were included in a backwards stepwise regression, metabolites not passing the cut-off threshold of *P*<0.05 were removed.



Figure 5-2 shows the metabolites significantly associated with intakes of each food group in the training dataset. Bars with a white centre represent the $-\log_{10} P$ -value from the discovery analysis. Solid bars indicate the $-\log_{10} P$ -value from the multivariate regression with the final metabolites included in the scores following data reduction. *Green* bars indicate a positive association and *red* bars indicate a negative association. Metabolites whose biochemical identity has not yet been identified are denoted by 'X-' followed by 5 digits.

Food group	Metabolite	Superpathway	Subpathway
Vegetables	↓ X-12063	Unknown	
Fruit	↑ stachydrine	Xenobiotics	Food component, Plant
	↑ threonate	Cofactors and vitamins	Ascorbate and aldarate metabolism
	↑ threitol	Carbohydrate	Nucleotide sugars, pentose metabolism
	↓ proline	Amino acid	Urea cycle; arginine-, proline-, metabolism
	↑ hippurate	Xenobiotics	Benzoate metabolism
Nuts and legumes	↑ tryptophan betaine	Amino acid	Tryptophan metabolism
Seafood	↑ 3-carboxy-4-methyl-5-propyl-2-furanpropanoate (CMPF)	Lipid	Fatty acid, dicarboxylate
	↓ 1-arachidonoylglycerophosphoethanolamine*	Lipid	Lysolipid
	↓ 1-oleoylglycerophosphoethanolamine	Lipid	Lysolipid
	↓ 1-eicosatrienoylglycerophosphocholine*	Lipid	Lysolipid
	↓ 1-linoleoylglycerophosphoethanolamine*	Lipid	Lysolipid
White meat	↑ 3-methylhistidine	Amino acid	Histidine metabolism
Red, processed meat and eggs	↑ trans-4-hydroxyproline	Amino acid	Urea cycle; arginine-, proline-, metabolism
	↑ creatine	Amino acid	Creatine metabolism
	↑ C-glycosyltryptophan*	Amino acid	Tryptophan metabolism
	↓ piperine	Xenobiotics	Food component, Plant
	↑ X-12696	Unknown	
Butter and cream	↑ nonanoylcarnitine*	Lipid	Carnitine metabolism
	↑ myristate (14:0)	Lipid	Long chain fatty acid
	↑ caprate (10:0)	Lipid	Medium chain fatty acid
	↑ X-02249	Unknown	
	↑ X-10510	Unknown	
Spreads and dressings	↑ X-11478	Unknown	
	↑ X-11521	Unknown	
	↑ X-11261	Unknown	

Table 5-5. List of metabolites associated with a single food group following the discovery analysis

Food group	Metabolite	Superpathway	Subpathway
Milk	↑ X-21365 (trimethyl-N-aminovalerate)	Unknown	
Теа	↑ N-acetylornithine	Amino acid	Urea cycle; arginine-, proline-, metabolism
Coffee	↑ catechol sulfate	Xenobiotics	Benzoate metabolism
	↑ paraxanthine	Xenobiotics	Xanthine metabolism
	↑ theophylline	Xenobiotics	Xanthine metabolism
	↑ X-05426	Unknown	
	↑ X-12217	Unknown	
Alcohol	↑ 4-androsten-3beta,17beta-diol disulfate 1*	Lipid	Sterol, Steroid
	↑ 2-oleoylglycerophosphocholine*	Lipid	Lysolipid
	↑ X-10395	Unknown	
	↑ X-11204	Unknown	
	↓ X-11444	Unknown	
	↓ X-11787	Unknown	

Table 5-5. List of metabolites associated with a single food group following the discovery analysis

5.3.2 Overall associations with food group intakes following data reduction

Each food group was associated with more than one metabolite (73 associations overall) which were potentially correlated, therefore a backward stepwise linear regression was performed to remove these using a significance threshold of P<0.05. **Table 5-4** summarises the results of this analysis, the detailed results can be found in **Appendix D Table 3**. For alcohol, seafood and coffee intakes, at least one metabolite was significant in the first model but in the opposite direction to the results of the fixed effects meta-analysis (**Appendix D Table 3**, metabolite name and result highlighted in red), therefore a second model was undertaken using only the significant metabolites that were in the same direction. The names and weightings (standardized betas) of the final metabolites used to produce the metabolite scores are highlighted in bold in **Appendix D Table 3**. The metabolites included in the final score are indicated by the solid bars in **Figure 5-2** which represent the $-\log_{10} P$ -value from the multivariate regression including all final metabolites in the food score (the R² for these regression models can be found in **Table 5-4**).

Complete results for each food group against each final metabolite score can be found in (**Appendix D Table 4**). The associations between the metabolite scores and food group intake were stronger than the top single metabolite association for all food groups in the training dataset. All top metabolites and score associations except for vegetables and spreads and dressings (continuous and weighted) passed the significance threshold for the discovery analysis (P<7.60x10⁻⁶) in the test group.

5.3.3 ROC analysis of metabolite food scores in the test group

Figure 5-3 shows the ROC curves for each of the metabolite scores ability to predict high and low consumers based on tertiles (intakes summarised in **Appendix D Table 5**). The results for the top metabolite and top performing metabolite score are presented in **Table 5-6** (results for all scores can be found in **Appendix D Table 6**). Overall, all scores were above the line of no-discrimination (AUC>0.5), indicating the metabolite scores performed better than a random guess at identifying low and high food group consumers. Metabolite scores for alcohol performed moderately (AUC>0.8) and metabolite scores for fruit, whole grains, seafood, fried foods, tea and coffee performed reasonably (AUC>0.7) at predicting high and low consumers. When comparing the combined metabolite scores against the top single metabolite from each score the weighted method performed significantly better (*P*<0.05) and the best for fruit,

seafood, red meat, sweet and sweet baked products, butter and cream and alcohol intakes. For the rest of the groups the scores did not significantly improve the prediction. For foods associated to only a single metabolite (fried foods, vegetables, nuts and legumes, and white meat), dividing the group into quartiles did not significantly change the predictive ability.

5.3.4 Metabolite score associations with food preferences in the test group

We previously showed that food preference patterns in our twin population are correlated to metabolite signatures in the blood, despite metabolomics profiles being measured seven years prior (Pallister et al., 2015). This finding suggests that food preference patterns are quite stable through time. Here I have used the food preference questionnaire to further validate the metabolite scores in the test group. All score associations passed the Bonferroni cut-off (P<3.33x10⁻³) except for spreads and dressings. The top results from this analysis can be found in **Table 5-7**. The full results are in **Appendix D Table 7**. For those foods only associated with 1 metabolite, vegetables and white meat were not significantly associated with food liking, which supports our results from the ROC analysis. These results provide further evidence that food liking has a strong impact on food intakes.

5.3.5 Metabolite scores results summary

- Summarising multiple metabolites into a score improved the strength of associations between food intakes and metabolite profiles.
- Differences in the overall predictive performance of scoring methods were negligible for most food groups, though the weighted method generally performed the best.
- iii) Based on the results from the ROC analysis and associations with food preferences, scores for vegetable, white meat and spreads and dressings did not appear useful.



Figure 5-3. ROC curves for each diet metabolite score

Each food group metabolite scoring method (quartiles, continuous and weighted) and the top associated metabolite were each fitted into a logistic regression model (adjusted for age, BMI, batch and family relatedness) to classify the top (1, positive outcome) and bottom (0, negative outcome) tertiles of food group intake. The equality of the receiver operating characteristic area (AUC) is shown for each scoring method and the top associated metabolite.



Figure 5-3. Continued...ROC curves for each diet metabolite score

Each food group metabolite scoring method (quartiles, continuous and weighted) and the top associated metabolite were each fitted into a logistic regression model (adjusted for age, BMI, batch and family relatedness) to classify the top (1, positive outcome) and bottom (0, negative outcome) tertiles of food group intake. The equality of the receiver operating characteristic area (AUC) is shown for each scoring method and the top associated metabolite.

Table 5-6. Results of ROC analysis ⁽¹⁾

						Versus top metabolite		
Food group	Score	Sensitivity	Specificity	Correctly classified	AUC	χ²	Р	
Vegetables ⁽²⁾	Top: X-12063	59.74%	55.47%	57.60%	0.6136 [0.5802, 0.6470]			
Eruit	Top: X-11315	65.59%	60.34%	62.98%	0.6783 [0.6479, 0.7087]			
Fluit	Weighted	66.84%	67.24%	67.04%	0.7411 [0.7131, 0.7690]	29.77	<0.0001	
Whole grains	Top: X-09789	65.25%	67.07%	66.15%	0.7134 [0.6841, 0.7428]			
whole grains	Weighted	67.63%	65.81%	66.72%	0.7224 [0.6934, 0.7514]	1.37	0.2425	
Nuts and legumes ⁽²⁾	Top: tryptophan betaine	64.25%	60.70%	62.47%	0.6723 [0.6366, 0.7080]			
Seafood	Top: DHA	64.79%	67.46%	66.13%	0.7230 [0.6943, 0.7516]			
Sealoou	Weighted	69.57%	70.68%	70.13%	0.7547 [0.7274, 0.7821]	11.88	0.0006	
White meat ⁽²⁾	Top: 3-methylhistidine	77.78%	32.44%	56.86%	0.5733 [0.5371, 0.6096]			
Pod most	Top: trans-4-hydroxyproline	58.94%	65.37%	62.16%	0.6697 [0.6390, 0.7003]			
Reumeat	Weighted	63.61%	65.53%	64.57%	0.6963 [0.6666, 0.7260]	9.36	0.0022	
Fried foods ⁽²⁾	Top: X-11372	65.30%	66.32%	65.81%	0.7322 [0.7038, 0.7606]			
Sweets and sweet baked products	Top: C-glycosyltryptophan*	58.67%	56.51%	57.59%	0.6002 [0.5679, 0.6325]			
Sweets and sweet baked products	Weighted	60.10%	58.97%	59.54%	0.6254 [0.5937, 0.6571]	5.07	0.0243	
Butter and cream	Top: nonanoylcarnitine*	71.88%	39.55%	56.84%	0.5889 [0.5529, 0.6249]			
	Weighted	59.15%	59.90%	59.52%	0.6385 [0.6036, 0.6734]	11.39	0.0007	
Spreads and dressings	Top: X-11261	59.08%	57.48%	58.28%	0.6199 [0.5880, 0.6518]			
	Quartiles	60.27%	60.10%	60.19%	0.6374 [0.6059, 0.6689]	5.42	0.0199	
Milk	Top: X-21365 [trimethyl-N-aminovalerate]	60.17%	60.65%	60.41%	0.6409 [0.6096, 0.6723]			
	Continuous	60.68%	61.33%	61.00%	0.6355 [0.6040, 0.6669]	0.22	0.6402	
Теа	Top: X-14473	59.28%	73.51%	67.04%	0.7384 [0.7078, 0.7691]			
	Quartiles	61.63%	64.63%	63.13%	0.7121 [0.6803, 0.7439]	2.68	0.1016	
Coffee	Top: X-14473	78.21%	65.38%	72.42%	0.7779 [0.7483, 0.8074]			
	Quartiles	72.06%	76.36%	74.21%	0.7917 [0.7627, 0.8208]	3.08	0.079	
Alcohol	Top: scyllo-inositol	75.75%	63.32%	70.02%	0.7531 [0.7230, 0.7832]			
Alcohor	Weighted	74.70%	76.40%	75.55%	0.8351 [0.8104, 0.8598]	38.99	<0.0001	

ROC, receiver operating characteristic; AUC area under the receiver operating characteristic curve

(1) Each food group metabolite scoring method (quartiles, continuous and weighted) and the top associated metabolite were each fitted into a logistic regression model (adjusted for age, BMI, batch and family relatedness) to classify the top (1, positive outcome) and bottom (0, negative outcome) tertiles of food group intake. The equality of the receiver operating characteristic area (AUC) for each scoring method was tested against the ROC area for the top associated metabolite.

(2) Only one metabolite marker

Food group	Top metabolite	Beta (SE)	Р	Best scoring method	Beta (SE)	Р
Vegetables ⁽²⁾	X-12063	-0.062 (0.059)	NS			
Fruit	X-11315	0.203 (0.045)	7.40x10 ⁻⁶	Weighted	0.077 (0.017)	1.12x10 ⁻⁵
Whole grains	X-09789	0.287 (0.054)	1.37x10 ⁻⁷	Weighted	0.045 (0.012)	1.70x10 ⁻⁴
Nuts and legumes ⁽²⁾	tryptophan betaine	0.271 (0.045)	3.04x10 ⁻⁹			
Seafood	DHA	0.333 (0.035)	6.85x10 ⁻²⁰	Weighted	0.173 (0.014)	1.51x10 ⁻³⁰
White meat ⁽²⁾	3-methylhistidine	0.083 (0.037)	2.45x10 ⁻² *			
Red meat	trans-4-hydroxyproline	0.221 (0.046)	2.29x10 ⁻⁶	Weighted	0.081 (0.009)	7.71x10 ⁻¹⁹
Fried foods ⁽²⁾	X-11372	0.344 (0.046)	2.69x10 ⁻¹³			
Sweets and sweet baked products	C-glycosyltryptophan	0.004 (0.002)	1.09x10 ⁻² *	Weighted	0.002 (0.000)	2.24x10 ⁻⁷
Butter and creams	nonanoylcarnitine	0.023 (0.033)	4.79x10 ⁻¹ *	Weighted	0.022 (0.007)	2.61x10 ⁻³
Spreads and dressings	X-11261	0.105 (0.041)	1.15x10 ⁻² *	Quartiles	0.207 (0.089)	2.06x10 ⁻² *
Milk	X-21365 [trimethyl-N-aminovalerate]	0.167 (0.041)	5.71x10⁻⁵	Continuous	0.352 (0.097)	3.15x10 ⁻⁴
Теа	X-14473	-0.217 (0.031)	7.83x10 ⁻¹²	Quartiles	0.624 (0.082)	8.88x10 ⁻¹⁴
Coffee	X-14473	0.241 (0.055)	1.36x10 ⁻⁵	Quartiles	0.689 (0.101)	1.98x10 ⁻¹¹
Alcohol	scyllo-inositol	0.322 (0.042)	1.01x10 ⁻¹³	Weighted	0.197 (0.018)	4.08x10 ⁻²⁷

Table 5-7. Top and metabolite score associations with food preferences in the test group ⁽¹⁾

NS, not significant; *, not statistically significant (P<3.33x10⁻³)
(1) The top metabolite associated with each food group and the best performing scoring method (if multiple metabolites associated) from the receiver operating characteristic curve analysis were using to predict preferences for food groups.
(2) Only one metabolite was associated with the food group.

5.3.6 Metabolite score associations with future incidence of MetS

Metabolite scores for butter and creams (OR[SE]: 0.602[0.119]; *P*=0.010), seafood (0.513[0.115]; *P*= 2.85×10^{-3}), and whole grains (0.325[0.095]; *P*= 1.15×10^{-4}) were associated with reduced odds of future MetS. Reported intakes for the same food groups were not significantly predictive of MetS, though effect directions were the same, which is suggestive that the metabolite scores were able to capture an effect more strongly than the questionnaires. Each significant metabolite score was predictive of at least one feature of MetS. Lower metabolite scores were predictive of reduced HDL-cholesterol for all food groups (**Table 5-8**). Lower metabolite scores for whole grains were mildly predictive of elevated triglycerides. There were no associations between metabolite scores and elevated WC, high blood pressure or impaired fasting plasma glucose.

Table 5-6. Metabolite Score predictions of enteria categories of the metabolic syndrome								
Criteria	Butter and creams		Seafo	od	Whole grains			
	OR[SE]	P	OR[SE]	P	OR[SE]	Р		
Elevated triglycerides		NS		NS	0.639[0.123]	0.020		
Reduced HDL- cholesterol	0.347[0.125]	3.38x10 ⁻³	0.413[0.123]	3.05x10 ⁻³	0.367[0.099]	2.12x10 ⁻⁴		

Table 5-8. Metabolite score predictions of criteria categories of the metabolic syndrome

NS, not significant (P<0.05); MetS, metabolic syndrome; HDL, high density lipoprotein cholesterol A logistic regression (0, no risk; 1, risk) was performed in 414 twins using each food metabolite score (residual adjusted for age, BMI and batch at time of metabolomics) to predict the 5 clinical MetS criteria taken \geq 5 years after metabolomics profiling.

5.4 Discussion

The primary aims of this chapter were to identify markers of long-term reported intakes of general food groups, to create biomarker scores using multiple methods and evaluate them, including testing them on the future development of MetS. Through the work I completed in this chapter I came to the following conclusions: i) summarising multiple metabolites into a score improved the strength of associations between food intakes and metabolite profiles; ii) differences in the overall predictive performance of scoring methods were negligible for most food groups, though the weighted method generally performed the best; iii) based on the results from the ROC analysis and associations with food preferences, I would not use scores for vegetable, white meat and spreads and dressings for further analysis. Nearly all of the metabolite associations identified and confirmed in discordant MZ twins have been identified in Chapter 4 as markers of reported intakes of more specific food items, I have therefore not highlighted these in the discussion. Moreover, I found that lower biomarker scores, though not reported intakes, associated to consumption of butter and creams, seafood and whole grain products associated to a higher risk of MetS later in life.

Few epidemiological studies and no twin studies to date have examined reported food intakes against metabolomics profiles to create biomarker scores and evaluated them. One study of African Americans examined the impact of alcohol consumption related metabolite subpathways on white blood cell levels and incident CVD (Zheng et al., 2014). In this chapter I used their method of summing quartile ranked metabolites to create a score, though this method did not perform the best overall and appears to lose some information. Moreover, while the authors created scores for different sub-pathways they did not acknowledge that these pathways are not specific to alcohol intake, such as 2-hydroxybutyrate and related branched chain amino acids which are markers of insulin resistance and type 2 diabetes (Menni et al., 2013a), and therefore may not truly connect alcohol consumption to CVD directly. I also used the summation score method that has previously been used to assess adherence to a healthy Nordic diet (Marklund et al., 2014). I found this method simple, though for most food groups did not perform better than the weighted method. I used the weighted method as it has been used to study markers of oxidative stress and their influence on incident, sporadic colorectal adenomas (Dash et al., 2013), though they did not directly test the difference in accuracy between the equal weight versus weighting methods, the authors did find the methods to predict colorectal adenomas

similarly. An issue with equally weighting scores is that they may be less biologically meaningful than weighting. However, the weights used here may not be applicable to other populations. Although through splitting our group into a training and test set I found the weighting method still tended to perform better. Single metabolites did not perform well compared with combining metabolites into scores, supporting the use of biomarker scores for future study.

All scores performed better than a random guess at identifying low and high food group consumers. Generally, scores including more metabolites performed better. Scores that performed with moderate accuracy (AUC>0.7) included those for alcohol, fruit, whole grains, seafood, fried foods, tea and coffee. Suggesting that these metabolites should be investigated more thoroughly. Previous metabolomics studies have identified strong associations with reported coffee, fruit, alcohol and seafood intakes (Guertin et al., 2014b, Guertin et al., 2015, Zheng et al., 2014), many of which I confirmed in this and the previous chapter. It is interesting, though encouraging, that the marker for fried food intake performed reasonably well as reporting of fried food intake may be more prone to underreporting. It is difficult to draw conclusions for metabolite scores that did not perform as well (AUC<0.7). However, the potential markers may simply not be good markers, though it also should be taken into account that ranking of the intakes for some foods may not be precise. To help resolve these issues controlled feeding studies should be undertaken, moreover applying different or new metabolomics platforms with more metabolites may yield markers with greater potential.

An important issue with creating scores is that many of the metabolites associated to food intakes appear to lack specificity to a particular food. Moreover, in our study many metabolites were associated to multiple foods that may also be a result of correlations in reporting and dietary patterns. I attempted to ameliorate this issue by running a multivariate linear regression including all foods associated to each applicable metabolite and used the stringent significance threshold from the discovery analysis. However, by doing this I may have eliminated important markers. Moreover, future studies may find that working with correlations in intakes and identifying markers of healthy versus unhealthy dietary patterns (which tend to cluster together) may be more useful for public health research (and easier for translation) than examining single foods alone.

I found that lower biomarker scores (though not reported intakes) for butter and creams, seafood and whole grains associated to a higher risk of MetS later in life. However, only 25 individuals subsequently developed MetS at a later time, therefore strong conclusions cannot be

drawn from this small sample size. It is interesting that metabolites associated to butter and cream intake were lower in individuals who later developed MetS, which seems conflicting. Moreover, higher scores on the butter and cream metabolite score were associated with higher levels of HDL-cholesterol. Although large population studies have supported these findings, a recent study on 15,105 Brazilian adults identified inverse associations between total dairy and high fat dairy consumption and later development of the MetS which appeared to be mediated by dairy saturated fat consumption (Drehmer et al., 2016). In a large population of adults from 2 US cohorts (the Health Professionals Follow-Up Study: 51,529 men; the Nurses' Health Study: 121,700 women), classic blood biomarkers of dairy fat consumption, trans palmitoleate, pentadecanoic and heptadecanoic acids were not associated with incidence of stroke (Yakoob et al., 2014). In another study of 659 adults from the triethnic multicenter Insulin Resistance Atherosclerosis Study, circulating pentadecanoic acid was inversely associated with incident type 2 diabetes, a clinical measure of insulin resistance and β -cell dysfunction (Santaren et al., 2014). I also found that individuals with lower whole grain metabolite scores were more likely to develop MetS. All of the metabolites forming the whole grain score are unknown therefore not much information can be derived from them at this time. Though it has been found that whole grain consumption appears to have a protective effect on MetS (Sahyoun et al., 2006) and cardiometabolic traits (Giacco et al., 2010). Using a biomarker approach, increased plasma alkylresorcinols (AR) ratio C17:0/C21:0, a biomarker of whole grain rye intake, was associated with an improved blood lipid profile following the consumption of a healthy Nordic diet in subjects with MetS (which includes high intakes of rye bread) (Magnusdottir et al., 2014), however total AR which is more representative of total whole grain wheat and rye intake was not associated with an improved lipid profile. I identified a particularly strong association between having lower scores on the whole grain metabolite score and having lower HDLcholesterol, though a strong relationship between HDL-cholesterol and whole grain consumption has not been identified (Hollaender et al., 2015). Higher scores on the seafood biomarker score were also found to associate with a reduced MetS risk and higher HDL cholesterol. A systematic review of 7 studies found 4 studies (one follow-up, three crosssectional) to suggest increased fish intake is protective of MetS (Torris et al., 2014). The long chain omega-3 fatty acids found in fatty fish, and also included in the biomarker score I used, are thought to be protective through improving vascular function and reducing inflammatory processes (Tousoulis et al., 2014). Supporting this, a recently published prospective study of

4356 American young adults found that increasing reported intakes of non-fried fish and dietary long-chain omega-3 fatty acids were associated inversely with MetS at 25 years follow-up (Kim et al., 2016). Biomarker studies are scarce on the relationship between blood long-chain omega-3 PUFAs and MetS risk. One study of chronic kidney disease patients found a pattern of high *n*-3 PUFAs in serum were not consistently associated with MetS (Huang et al., 2014). In another study serum DHA was associated with reduced cardiometabolic risk (Song et al., 2014).

There are a number of limitations and factors that must be considered. The Metabolon method does not determine the true blood concentration of the metabolites therefore I could not identify accurate cut-off points for what constitutes high and low intakes or risk levels. For chocolate, refined grains, fermented dairy, soy and other milk, and soda intakes no metabolite biomarkers were identified or did not pass data reduction and could not be studied further. By evaluating the metabolite scores against FFQ data I likely did not capture the true discriminatory ability of the scores due to the nature of dietary self-reporting. Through using these methods I was able to compare the utility of different scoring methods. Future feeding studies will be better equipped to assess biomarker utilization more accurately. The classification of foods was done a priori based on foods derived from similar sources (e.g. plants, animals, processing), which confer similar health benefits (e.g. whole grain versus refined grain), food usage and cooking. I may have chosen foods to be in a group that are not well correlated to intakes of the rest of the foods, therefore one food may be more strongly related to a metabolite, whereas another food included in the group has no relationship. Future work should aim to unravel the precise mechanism for these associations in order to discern which foods contribute to the same metabolite associations. Only women were used for this study, therefore the results may not be applicable to men. Many markers are unidentified at this time therefore I could not confirm they may be biologically related to food intakes, however I used stringent cut-offs, discordant MZ twins and data reduction methods to identify the best metabolite markers possible in our population. Importantly, the metabolite scores were tested in a group independent from the discovery group.

Chapter 6 Metabolomic associates of microbial diversity, their modulation by diet and relationship to the metabolic syndrome

In this chapter I analysed gut microbiome Shannon diversity (Shannon) for its association with blood metabolites from the Metabolon platform, I then examined whether these microbe-related metabolites were associated with reported food intakes and finally, I explored longitudinal changes of the top diversity-associated metabolite and its association with metabolic syndrome risk and its components.

6.1 Introduction

The microbiome refers to the collective genomes of the microorganisms within an environmental niche. Within the human gut, microbiota undertake important processes allowing for potential host-microbe interactions such as vitamin synthesis, hormone production, metabolism of food components and interactions with host innate immunity. The number of different organisms present within the human gut (referred to as 'richness' or 'diversity') is associated with the increased abundance of beneficial bacteria and is emerging as an important indicator of health. Richness represents the total gene count, while diversity additionally accounts for the rarity of those microbial genes present within the whole dataset. Reduced alpha-diversity (intraindividual diversity) has been coupled with dysbiosis (microbial imbalance) in inflammatory bowel disease patients (Manichanh et al., 2006, Michail et al., 2012, Scher et al., 2015), though this has not been highlighted in studies of obesity (Turnbaugh et al., 2009) or metabolic health (Le Chatelier et al., 2013). Although mechanisms are unclear microbial imbalance occurs when beneficial bacterial growth is compromised and a few pathogenic taxa dominate and overtake the metabolic potential of the microbiome with possible deleterious effects. Recent studies have shown reduced diversity is present in disease states and metabolic conditions such as autoimmune diseases, including inflammatory bowel disease (Michail et al., 2012, Ott et al., 2004, Manichanh et al., 2006) and in patients with psoriatic arthritis and inflammatory bowel disease (Scher et al., 2015), and metabolic derangements, including obesity (Turnbaugh et al., 2009) and MetS-related phenotypes (Le Chatelier et al., 2013). Diet has been shown to contribute to gut microbiome richness (Cotillard et al., 2013), in particular dietary fibre promotes stability in richness in humans (Tap et al., 2015) and this stability has recently been shown to be transferred to future generations in mice (Sonnenburg et al., 2016), highlighting the important potential of good nutrition in humans.

Microbes play a distinct role in human metabolism by transforming food- and hostderived metabolites, such as bile acids, polyphenols and fibre. These products may in turn influence disease development by interacting with human host physiology. Studies are now attempting to untangle these interactions through measuring the chemical profile (metabolome) in biofluids (urine and blood), faeces, tissues and organs in collaboration with microbiome analysis. Comparing conventional versus germ-free mice, conventional mice exhibited elevated blood levels of indole-containing compounds (e.g. indoxyl sulfate and indole-3-propionic acid), serotonin, sulfated compounds (e.g. phenyl and p-cresol sulfate), and glycine-conjugated compounds (hippuric acid, cinnamoylglycine and phenylpropionylglycine), showing the major contribution of the gut microbiome to metabolism (Wikoff et al., 2009). In another study, urinary excretion of hippuric acid, a metabolite derived from high polyphenol foods (Gonthier et al., 2003, Walsh et al., 2007), has been found to discriminate well between obese and normal weight controls (Calvani et al., 2010). Many of the metabolites implicated are food-derived components. In the two previous chapters I found many food associations with microbial cometabolites, therefore merging microbiome and metabolomics approaches with dietary studies is the logical next step in improving our understanding of the interplay between diet, the microbiome and metabolic disease.

Diet is thought to be the most important avenue for modulating the gut microbial composition and its metabolic outputs, despite this there are relatively few good human studies. It has been shown that daily consumption of 40 g of dark chocolate for 2 weeks altered urinary output of gut microbial metabolites, increasing hippurate and methylamines, and reducing pcresol sulfate (Martin et al., 2009). A recent randomized controlled pilot found consuming 30 g/d of heat-stabilized rice bran for 28 days increased abundance of 8 operational taxonomic units Methanobrevibacter, Paraprevotella, Ruminococcus, (OTUs) from the genera Dialister, Anaerostipes and Barnesiella, and 3 from Bifidobacterium and Ruminococcus, moreover elevated levels of secondary bile acids (deoxycholic and lithocholic) and metabolites derived from plants and microbial modifications of plant phenolics (benzoic, hydrocinnamic and phosphoric acids, and inositol monophosphate) were observed in the faecal metabolome (Sheflin et al., 2015). The microbial metabolism of choline and L-carnitine, mainly found in meat,

fish and eggs, to trimethylamine (TMA) and then oxidized in the liver to trimethylamine-N-oxide (TMAO) has been implicated in why some red-meat consumers develop atherosclerosis (Koeth et al., 2013, Wang et al., 2011). Specifically, TMAO has been suggested to encourage the upregulation of macrophage scavenger receptors and through this enhance forward cholesterol transport (Bremer, 1983). Though in mice with intact intestinal microbiota and also with higher TMA and TMAO blood concentrations, choline and L-carnitine feeding each suppressed reverse cholesterol transport (Koeth et al., 2013). Interestingly, suppression of the gut microbiota removed the effect.

The metabolomic signature of gut microbial alpha-diversity has yet to be examined on a large scale, and could provide useful candidate markers of a metabolically fit microbiome. Therefore for the current chapter, I aimed to:

- i) Identify the blood metabolites correlated with gut microbiome diversity.
- ii) Examine the impact of food intake on these metabolites.
- Examine if longitudinal changes in these metabolites are predictive of both future metabolic syndrome and its components.
6.2 Materials and methods

Data relevant to this chapter included the reported intakes of 20 food groups as described in the previous chapter (**Section 5.3.1**), microbiome data and blood metabolomics data (details **Section 3.1.6.1**). I used all 292 known metabolites measured by the Metabolon platform. Twins who completed FFQs collected between 1995 and 2001 and in 2007 were used in the discovery analysis (*n*=1529). New FFQ data were collected between 2014 and 2015, 484 additional twins had microbiome, metabolomics data and completed FFQs during this time, 420 of these twins who had no co-twin in the discovery sample were used as a validation sample. The remaining 64 twins who had co-twins in the discovery sample were used for the subsequent microbiome and MetS analyses.

 Table 6.1 provides the study population characteristics and subject numbers.

	Whole	FFQ<2014	FFQ≥2014
		(Discovery)	(including validation)
	Mean (SD)	Mean (SD)	Mean (SD)
Ν	2013	1529	484
Age (y)	57.2 (10.6)	57.7 (10.6)	55.4 (10.4)
BMI	26.0 (4.6)	26.1 (4.6)	25.9 (4.6)
Sex (M:F)	113:1909	0:1535	113:374
MZ:DZ pairs	408:392		
Singletons	421		
Food groups			
(servings/week)			
Vegetables	34.9 (16.5)	34.8 (15.4)	35.1 (19.6)
Fruit	21.4 (12.7)	21.9 (12.4)	19.5 (13.5)
Whole grains	10.3 (8.0)	10.7 (8.1)	9.2 (7.4)
Refined grains	8.6 (7.6)	9.0 (7.9)	7.2 (6.3)
Nuts and legumes	7.9 (5.8)	7.6 (5.2)	8.8 (7.2)
Seafood	2.4 (2.0)	2.5 (2.0)	2.3 (2.1)
White meat	1.9 (1.3)	1.9 (1.3)	1.9 (1.3)
Red meat	6.9 (4.1)	6.8 (3.9)	7.4 (4.6)
Fermented dairy	6.2 (5.1)	6.1 (4.7)	6.5 (6.2)
Fried foods	4.5 (3.5)	4.5 (3.3)	4.8 (4.1)
Sweets and sweet	15.2 (13.9)	15.5 (14.0)	14.1 (13.4)
baked products			
Chocolate	4.0 (5.9)	4.0 (5.8)	3.9 (6.3)
Butter and cream	4.4 (6.5)	4.2 (6.4)	5.0 (6.8)
Spreads and	8.2 (9.0)	8.5 (9.1)	7.2 (8.6)
dressings			
Milk	3.2 (2.3)	3.4 (2.3)	2.8 (2.2)
Soya and other milk	0.2 (0.8)	0.2 (0.9)	0.2 (0.8)
Soda	1.7 (4.2)	1.8 (4.3)	1.3 (3.5)
Tea	19.3 (13.5)	19.2 (13.5)	19.4 (13.5)
Coffee	9.0 (10.6)	9.1 (10.7)	8.8 (10.5)
Alcohol	6.1 (8.0)	6.2 (7.9)	5.9 (8.4)

Table 6-1.	Study	population	characteristics	for	the	whole,	discovery	and	validation
samples									

6.2.1 Faecal microbiome composition

Faecal samples were collected at follow-up and the composition of the gut microbiome was determined by 16S rRNA gene sequencing carried out at Cornell as previously described (Goodrich et al., 2016). Firstly, the V4 region of the 16S rRNA gene was amplified and then sequenced on Illumina MiSeq. The reads were next compiled to operational taxonomic units (OTUs) (Ley et al., 2006). Quality control was undertaken by Matthew Jackson and Tiphaine Martin at KCL by sample, paired-ends with an overlap of less than 200nt were removed. Chimeric sequences were then removed by de novo chimera detection in USEARCH (Edgar et al., 2011). Using Sumaclust within QIIME 1.9.0 de novo OTU clustering was undertaken across all reads, reads with a 97% identity threshold were brought together (Jackson et al., 2016a, Caporaso et al., 2010). Log transformation was undertaken on OTU counts, prior to transformation zero counts were given an arbitrary value (1x10⁻⁶). OTU abundances were then adjusted for technical covariates including sequencing depth, sequencing run, sequencing technician and sample collection method using linear modelling and the residuals obtained. As the residuals were not normally distributed, I completed an inverse normalisation. To determine alpha diversity, the total OTU count table was rarefied to 10000 sequences for each sample 50 times. Per sample, alpha diversity metrics were determined in each of the rarefied tables and the average score for all 50 was considered as the final diversity measure. The primary alpha diversity metric considered was the Shannon Index as this was considered the most robust and is a commonly used metric though findings were confirmed on other parameters including observed OTU counts and Chao1 (richness), and the Simpson diversity index. All alpha diversity indices were standardised to have mean 0 and SD 1.

6.2.2 Classification of the metabolic syndrome

I determined MetS status using the criteria outlined by the International Diabetes Federation and the American Heart Association/National Heart, Lung, and Blood Institute (Alberti et al., 2009). Consult **Section 5.2.4** for details.

Table 6-2 provides the clinical characteristics of the subsample of twins studied.

	Londitudinal sample (n=1032)
	Mean (SD)
Age at MetS status (y)	64.2 (7.8)
MetS status (0, no; 1, yes)	906:116
Longitudinal metabolomics baseline to endpoint (y)	10.6 (3.9)
Sex (M:F)	27:1005
BMI (kg/m2)	26.2 (4.5)
Systolic blood pressure (mmHg)	129.0 (15.7)
Diastolic blood pressure (mmHg)	75.5 (9.7)
Glucose (mmol/L)	4.9 (0.6)
Cholesterol (mmol/L)	5.6 (1.0)
HDL-Cholesterol (mmol/L)	1.9 (0.5)
Triglycerides (mmol/L)	1.1 (0.5)

Table 6-2. Clinical characteristics of the twin subsample to investigate longitudinal hippurate, diversity, diet with the MetS phenotypes and its components

6.2.3 Statistical analysis

Statistical analysis was carried out using Stata version 12. The statistical analysis was undertaken in two parts. First, a marker of microbiome diversity was identified, its relationship to food intake explored and associations with microbiome OTUs/collapsed taxonomies identified. In the second part, the relationship between longitudinal levels of the diversity metabolite marker, diet, diversity and associated OTUs/taxa with the risk of MetS and its components were explored.

6.2.3.1 Part 1: Metabolite associations with diversity, relationship to food intake and associations with microbiome OTUs/taxa

Figure 6-1 shows the analysis pipeline for part 1.

6.2.3.1.1 Microbiome diversity-metabolite associations

I ran a linear regression model with Shannon diversity as a predictor of the metabolite level (292 metabolites, Metabolon platform) in a group of 1529 female twins, adjusted for age, batch, BMI and family relatedness (closest to blood sampling) and multiple testing (Bonferroni: $0.05/292 = 1.71 \times 10^{-4}$). Significant metabolites from the discovery sample were then evaluated against Shannon diversity using the same linear regression (additionally including sex as a covariate) in the male (*n*=113) and female validation sample (*n*=420), associations passing the 5% level of significance were considered validated.

6.2.3.1.2 Food intakes associated with diversity-associated metabolites

To identify if any validated diversity-associated metabolites were potentially related to food intake, in the discovery sample I used reported intakes of the 20 food groups to predict levels of the metabolites (Bonferroni: $0.05/[5 \text{ metabolites } x 20 \text{ food groups}] = 5.00 \times 10^{-4}$), adjusted for the same covariates as above. In a subsample of individuals from the discovery group (*n*=788) longitudinal metabolomics data were available (*n*=705 3 time points, *n*=83 2 time points) as well as reported food group intake at the same time or 5 years before the first blood sample.



I determined trajectories of change in the top diversity-associated metabolite (years after baseline: 2^{nd} time point: 7.4 [range: 2.3-12.8]; 3^{rd} time point: 13.9 [range: 8.3-17.9]) by empirical Bayes predictions (adjusted for age and BMI) which estimates the rate of change in standard deviations/year (Rabe-Hesketh and Skrondal, 2008). Using this method point estimates were calculated and a slope of change determined. Food group intake at baseline was then used to predict the metabolite trajectories (Bonferroni: 0.05/20 food groups = 0.0025).

A predictive score was created from those significantly associated foods according to the direction of association (i.e. positive association: Q1=0, Q2=1, Q3=2, Q4=3; negative association: Q1=3, Q2=2, Q3=1, Q4=0) and summed. In the validation sample, I conducted a logistic regression model (adjusted for covariates, including sex) to test the ability of the top diversity-associated metabolite to identify twins in the top and bottom tertiles of Shannon diversity, as well as the utility of the diet score to identify the top and bottom tertiles of the metabolite in blood.

6.2.3.1.3 Food-microbiome-metabolite axis

To identify associations between the metabolite and the microbiome I combined the discovery and validation samples (n=2013). To establish the strong association between the metabolite and diet score with richness (observed OTUs) and additional diversity metrics (Simpson and Chao1), I ran these associations using the same linear regression as for the Shannon diversity discovery analysis in the pooled sample. I then evaluated associations within the microbiome by running a linear regression model using the OTUs and OTUs collapsed at each taxonomic level (phylum, class, order, family, genus) as predictors of the metabolite adjusted for covariates, Shannon diversity and multiple-testing (Bonferroni cut-off; Table 6-3 shows the significance threshold for the OTUs and each taxonomic level). To determine the total variance in both Shannon diversity and the metabolite accounted for by metabolite-associated OTUs, I included all associated OTUs in a backwards stepwise linear regression using P<0.05 as the threshold cut-off, I report the R² for each model. Metabolite-associated OTUs/taxa were examined for their association with the diet score (predictor) adjusted for covariates, the metabolite and multiple testing (assigned at each taxonomic level; **Table 6-3**). To investigate if any of the foods forming the score were driving associations, I ran a multivariate regression model including all metabolite-associated foods and the same covariates.

	Hippurate		Hippurate diet ⁽²⁾			
Level	Number of variables	Р	Number of variables	Р		
Phylum	34	1.47x10 ⁻³	1	0.05		
Class	72	6.94x10 ⁻⁴	2	0.025		
Order	137	3.65x10 ⁻⁴	3	0.017		
Family	194	2.58x10 ⁻⁴	3	0.017		
Genus	382	1.31x10 ⁻⁴	7	7.14x10 ⁻³		
OTU	581	8.61x10 ⁻⁵	30	1.67x10 ⁻³		

Table 6-3. Statistical significance thresholds for microbiome analysis using Bonferroni correction ⁽¹⁾

(1) Bonferroni correction was calculated within each level.

(2) Variables significantly associated with hippurate were tested for their association against the hippurate diet score.

6.2.3.1.4 Microbiome-metabolite-diet interactions

I next tested if there was an interaction between the diet score and the microbiome OTU/collapsed taxa associated to both the metabolite and the diet score in the prediction of the blood metabolite levels. I first ran a continuous x continuous interaction model using each OTU/collapsed taxa adjusting for covariates (age, BMI, batch, sex and Shannon diversity). In the second analysis, to determine if total numbers of all associated bacteria were important, I first multiplied abundances of associated microbiome variables by the direction of association with the metabolite (-1, negative, 1, positive) and summed the variables. I then ran the same continuous x continuous interaction model as above using this variable.

6.2.3.2 Part 2: The relationship between longitudinal levels of the diversity metabolite marker, diet, diversity and associated OTUs/taxa with the risk of MetS and its components

Figure 6-2 shows the analysis pipeline for part 2.

6.2.3.2.1 Relationship between microbiome diversity, longitudinal metabolite and the MetS score and its components

Microbiome diversity has been reduced in metabolic diseases, therefore I hypothesized that reduced longitudinal changes in the top metabolite diversity marker would be associated with increased MetS risk. A subsample of 1032 individuals had longitudinal blood metabolite levels (n=533 with 2 time points, n=499 with 3 time points). The time for the longitudinal analysis ranged from 2.4-17.9 years. I evaluated whether the longitudinal levels of the top diversity

metabolite predicted MetS status and scoring or any of its components. I ran a linear regression model using Shannon diversity, the metabolite trajectory, the diet score or metabolite/diet-associated OTUs/taxa to predict MetS status (adjusting for age, sex, and family relatedness), and each component adjusting for age, BMI (except for BMI), sex, and family relatedness.

6.2.3.2.2 Metabolite trajectory association with MetS (and components) mediated by Shannon diversity, the diet score and specific OTUs/taxa

The aim of this analysis was to determine the degree to which the variance in MetS (and components) attributable to the metabolite trajectory was accounted for by the MetS (and components) association with Shannon diversity, the diet score or specific OTUs/taxa. The proportion of the variance of the MetS status and its components were determined for the metabolite trajectory after taking into account all covariates. This quantity is indicated as r_x^2 . The proportion of the variance for the MetS status (and its components) explained by the metabolite trajectory was then calculated after taking into account the same covariates as above but also including, Shannon diversity, the diet score (if applicable) or associated OTUs/taxa (r_{xy}^2). The percentage of the longitudinal metabolite association mediated by Shannon diversity, the diet status (and components) that is due to the longitudinal metabolite trajectory association with Shannon diversity, the diet score and associated taxa, namely $1 - (r_{xy}^2/r_x^2)$.

6.2.3.2.1 Confirmation of findings in twins discordant for Shannon diversity

Fifty-five MZ twin pairs discordant (\geq 1 SD) for Shannon diversity were used to confirm associations with diversity associations with the top metabolite cross-sectionally and longitudinally, the diet score and significant MetS components.



6.3 Part 1 results

6.3.1 Microbiome diversity metabolomics associations

Eight metabolites significantly correlated with Shannon diversity in the discovery sample and five were validated in the validation sample (**Table 6-4**). Among those, hippurate, a benzoate metabolite, associated significantly (P<5.00x10⁻⁴: 0.05/[5 metabolites x 20 food groups]) with intakes of fruit (0.012[0.002]; P=7.36x10⁻⁸) and whole grains (0.013[0.003]; P=2.05x10⁻⁵). Another benzoate metabolite 3-phenylpropionate was associated with fried foods (-0.045[0.009]; P=5.63x10⁻⁷), whole grains (0.018[0.004]; P=2.71x10⁻⁶) and fruit (0.010[0.002]; P=2.45x10⁻⁵). Hippurate and 3-phenylpropionate were strongly correlated (r=0.51), although summing the two metabolites did not improve their association with Shannon diversity (hippurate R²: 0.0258; 3-phenylpropionate R²: 0.0122; and combined R²: 0.0236), therefore for the remainder of the analysis I focused on hippurate.

6.3.2 Food intakes predict longitudinal hippurate trajectories

I next analysed longitudinal levels of hippurate (**Figure 6-3** shows the data for 25 randomly selected twins). Higher intakes of whole grains, coffee and fruit significantly (*P*<0.0025) predicted increasing hippurate trajectories (**Table 6-5**). All associations remained significant in a multivariate linear regression (**Table 6-5**) and together accounted for 5.3% of the variance in the hippurate trajectory. The nutrient profile of the score formed from these foods is shown in **Table 6-6**. I validated the hippurate diet score in the validation sample against hippurate (0.089[0.024]; $P=3.21\times10^{-4}$) and Shannon diversity (0.040[0.019]; P=0.035), independently of hippurate. The diet score was moderately ($h^2 \ge 30\%$) heritable (A: 0.3782 [0.3024, 0.4485]; E: 0.6218 [0.5515, 0.6976]), with the AE model being the best fit.



Table 6-4. Metabolites associated to Shannon diversity in the discovery sample, following backward stepwise linear regression and in the validation sample ⁽¹⁾

			Discovery (<i>n</i> =1529)		Validation (<i>n</i> =420) ⁽²⁾	
Metabolite	Super- pathway	Sub-pathway	beta (SE)	P	beta (SE)	Р
Hippurate	Xenobiotics	Benzoate metabolism	0.230 (0.040)	3.72x10 ⁻⁸	0.238 (0.072)	0.001*
p-cresol sulfate	Amino acid	Phenylalanine & tyrosine metabolism	0.200 (0.040)	9.90x10 ⁻⁸	0.179 (0.063)	0.005*
phenol sulfate	Amino acid	Phenylalanine & tyrosine metabolism	-0.200 (0.040)	5.82x10 ⁻⁷	-0.121 (0.063)	0.055
Phenylacetylglutamine	Amino acid	Phenylalanine & tyrosine metabolism	0.180 (0.040)	5.21x10 ⁻⁶	0.195 (0.062)	0.002*
3-phenylpropionate (hydrocinnamate)	Amino acid	Phenylalanine & tyrosine metabolism	0.160 (0.040)	3.43x10 ⁻⁵	0.185 (0.084)	0.028*
4-ethylphenylsulfate	Xenobiotics	Benzoate metabolism	0.190 (0.050)	5.12x10 ⁻⁵	0.062 (0.081)	0.441
Hyodeoxycholate	Lipid	Bile acid metabolism	-0.190 (0.050)	8.66x10 ⁻⁵	-0.215 (0.089)	0.016*
Indolepropionate	Amino acid	Tryptophan metabolism	0.140 (0.040)	9.20x10 ⁻⁵	0.093 (0.083)	0.262

*Statistically significant: P<0.05

(1) A linear regression was performed using Shannon diversity to predict levels of 292 metabolites adjusting for age, BMI, batch effects (and sex in the validation) and family relatedness.

(2) Statistically significant ($P < 1.71 \times 10^{-4}$) associations from the discovery group were validated in the validation group.

Table 6-5. Food intake and score predictions of longitudinal hippurate trajectories and following a backwards stepwise linear regression in a subsample of 788 twins

	Independent regression ⁽¹⁾	Multivariate regression ⁽²⁾
	Beta (SE) P	Beta (SE) P
Whole grains	1.70x10 ⁻⁴ (3.84x10 ⁻⁵) 9.54x10 ⁻⁶	1.58x10 ⁻⁴ (3.83x10 ⁻⁵) 3.90x10 ⁻⁵
Coffee	$1.03 \times 10^{-4} (2.82 \times 10^{-5}) 2.73 \times 10^{-4}$	1.14x10 ⁻⁴ (2.79x10 ⁻⁵) 4.46x10 ⁻⁵
Fruit	8.43x10 ⁻⁵ (2.71x10 ⁻⁵) 1.89x10 ⁻³	$7.90 \times 10^{-5} (2.70 \times 10^{-5}) = 3.48 \times 10^{-3}$
Diet score	1.07x10 ⁻³ (1.71x10 ⁻⁴) 7.18x10 ⁻¹⁰	

(1) Intakes of 20 food groups at baseline were used to predict the hippurate trajectory adjusted for age, BMI and sex. Statistical significance was defined as P<0.0025 (0.05/20 food groups). The diet score was formed by summing quartile ranked intakes of food groups significantly associated with the hippurate trajectory.</p>

(2) Whole grain, coffee and fruit intake were included in a multivariate regression to predict the hippurate trajectory including age, BMI and sex as covariates.

Table 6-6. Nutrient profile and linear trends of the hippurate diet score according to tertiles of the hippurate diet score

	Tertile 1	Tertile 2	Tertile 3	Trend	l
Nutrient	Mean (SD)	Mean (SD)	Mean (SD)	Beta (SE)	P
Energy (kcal)	2009.4 (562.8)	1847.1 (534.2)	1826.6 (498.7)	-90.77 (14.75)	9.16x10 ⁻¹⁰
Fat (g/d)	72.2 (12.5)	67.2 (10.4)	62.5 (9.9)	-4.87 (0.3)	9.54x10 ⁻⁵⁵
Saturated FAs (g/d)	26.8 (6.6)	23.7 (5.8)	21.2 (5.2)	-2.76 (0.16)	8.00x10 ⁻⁶¹
MUFAs (g/d)	24.2 (4.6)	22.1 (3.8)	20.2 (3.7)	-1.98 (0.11)	2.19x10 ⁻⁶⁴
PUFAs (g/d)	15.6 (5.2)	15.8 (4.3)	15.7 (4)	0.04 (0.13)	NS
trans-FAs (g/d)	1.8 (0.7)	1.5 (0.6)	1.3 (0.5)	-0.22 (0.02)	2.47x10 ⁻³⁷
Cholesterol (mg/d)	241.4 (92.2)	233.1 (78.5)	210.8 (70.7)	-15.34 (2.24)	8.98x10 ⁻¹²
Protein (g/d)	79.3 (13)	80.5 (12.7)	80.4 (11.5)	0.56 (0.34)	NS
Carbohydrate (g/d)	221.6 (39.3)	232.2 (32)	245.2 (29.8)	11.83 (0.93)	2.14x10 ⁻³⁵
Starch (g/d)	112.3 (31.2)	108.8 (26.6)	111.2 (27)	-0.54 (0.78)	NS
Total sugars (g/d)	106.6 (30.6)	120.7 (28.2)	131.1 (26.2)	12.23 (0.78)	8.44x10 ⁻⁵²
Glucose (g/d)	17.8 (7.8)	23.4 (9.1)	27.1 (7.9)	4.64 (0.23)	2.42x10 ⁻⁸²
Fructose (g/d)	19.8 (9.1)	27.5 (11.2)	32.9 (9.8)	6.57 (0.28)	1.02×10^{-107}
Sucrose (g/d)	44.3 (18.9)	44.2 (14.2)	45.5 (12.8)	0.58 (0.43)	NS
Maltose (g/d)	3.4 (1.9)	3.1 (1.5)	3.3 (1.5)	-0.03 (0.05)	NS
Lactose (g/d)	18.2 (10.7)	17.7 (9.8)	17.3 (9.2)	-0.45 (0.27)	9.79x10 ⁻²
NSP (g/d)	16.8 (5)	20.4 (4.9)	23.4 (5.2)	3.29 (0.14)	9.35x10 ⁻¹¹⁰
Alcohol (g/d)	10.6 (14.8)	9.6 (11.9)	8.6 (10.5)	-0.95 (0.35)	5.77x10 ⁻³
Water (g/d)	2395.3 (596.3)	2623.5 (575.6)	2817.4 (591.6)	210.9 (16.26)	5.21x10 ⁻³⁷
Sodium (mg/d)	2172.9 (487.6)	2263 (502.5)	2341.4 (468.7)	84.19 (13.47)	4.98x10 ⁻¹⁰
Potassium (mg/d)	3526.7 (575.9)	3886 (576.2)	4174.1 (564.8)	323.4 (15.84)	2.50x10 ⁻⁸⁴
Chloride (mg/d)	3446.9 (750.3)	3603.9 (773)	3757.3 (712.8)	155.16 (20.65)	8.56x10 ⁻¹⁴
Calcium (mg/d)	1015.3 (288)	1037.2 (276.3)	1037.9 (274.2)	11.17 (7.73)	NS
Magnesium (mg/d)	299.1 (48.6)	343.4 (48.3)	377.1 (50.4)	38.93 (1.36)	2.03×10^{-151}
Phosphorous (mg/d)	1388.4 (223.7)	1476.2 (213.8)	1528.9 (207.9)	70.11 (5.96)	5.50x10 ⁻³¹
Iron (mg/d)	11.5 (2.5)	12.8 (2.7)	13.7 (3)	1.08 (0.08)	1.64x10 ⁻⁴³
Copper (mg/d)	1.5 (0.5)	1.6 (0.5)	1.6 (0.4)	0.08 (0.01)	5.57x10 ⁻⁹
Zinc (mg/d)	9.8 (1.7)	10.2 (1.6)	10.3 (1.5)	0.27 (0.04)	2.00x10 ⁻⁹
Manganese (mg/d)	3.7 (1.1)	4.1 (1.1)	4.5 (1.1)	0.41 (0.03)	2.02x10 ⁻³⁹
Iodine (ug/d)	199.3 (74.8)	205.7 (72.6)	206.1 (67.2)	3.35 (1.98)	9.12x10 ⁻²
Retinol (ug/d)	610 (776.9)	500.6 (508.6)	439.2 (425.7)	-85.14 (16.18)	1.58x10 ⁻⁷
Carotene (ug/d)	4987.8 (3370.1)	5754.6 (3645.7)	5896.5 (3627.5)	451.48 (98.45)	4.80x10 ⁻⁶
Vitamin D (ug/d)	2.3 (1)	2.5 (1)	2.5 (1.2)	0.1 (0.03)	5.47x10 ⁻⁴
Vitamin E (mg/d)	10.1 (3.2)	11.2 (3.2)	11.7 (3.1)	0.78 (0.09)	9.48x10 ⁻¹²
Thiamin (mg/d)	1.6 (0.4)	1.7 (0.4)	1.8 (0.4)	0.1 (0.01)	1.15x10 ⁻²¹
Riboflavin (mg/d)	2.2 (0.7)	2.2 (0.7)	2.3 (0.6)	0.02 (0.02)	NS
Niacin (mg/d)	19.8 (4.9)	21.2 (4.7)	23 (4.6)	1.6 (0.13)	2.67x10 ⁻³³
Tryptophan (mg/d)	16.6 (2.7)	16.8 (2.6)	16.9 (2.3)	0.17 (0.07)	1.59x10 ⁻²
vitamin B6 (mg/d)	2.3 (0.6)	2.5 (0.6)	2.7 (0.5)	0.16 (0.02)	0.41×10^{-20}
Vitamin B12 (mg/d)	6.2 (3)	6.1 (2.4)	5.9 (2.2)	-0.16 (0.07)	1.87x10 ⁻²
Folate (ug/d)	358 (106.9)	388.7 (113.4)	416 (110.1)	28.97 (3.05)	6.00×10^{-21}
Pantothenate (mg/d)	5.7 (1.6)	0.2 (5.3)	0.2 (1.8)	0.24 (0.09)	1.11X10
Biotin (ug/d)	40.9 (9.8)	45.4 (9.1)	49.4 (8.9)	4.25 (0.26)	4.40x10 ⁻⁵⁶
Vitamin C (mg/d)	126.8 (58.5)	169.6 (72.2)	200.6 (74.1)	36.88 (1.91)	1.37×10^{-70}

NS= not significant: P>0.05. FA: fatty acid, MUFAS: monounsaturated fatty acids, PUFAS:

polyunsaturated fatty acids, NSP: non-starch polysaccharides Tertile 1: score 0-3; tertile 2: score 4-5; tertile 3: scores 6-9. Linear trend determined by using the tertile of the hippurate diet score as a predictor of the energy-adjusted nutrient intake.

6.3.3 Utility of hippurate as a marker of diversity and the hippurate diet score as

a marker of hippurate

Overall hippurate measured at a single time point correctly classified 64.64% of individuals into the upper and lower tertiles of Shannon diversity (sensitivity: 68.57%; specificity: 60.71%). The AUC was 0.682 (0.620, 0.745) (**Figure 6-4**). The hippurate diet score correctly classified 60.93% of individuals into the upper and lower tertiles of hippurate levels (sensitivity: 64.03%; specificity: 57.86%). The AUC was 0.649 (0.585, 0.713) (**Figure 6-5**). This suggests both models had low accuracy.



6.3.4 Associations with hippurate and the hippurate diet score across diversity

metrics

I confirmed the same directional effects for other diversity metrics for hippurate and the diet

score (Table 6-7).

Table 6-7. Associations between hippurate and the hippurate diet score across diversity metrics in the whole sample (3)

	Hippurate **/		Hippurate die	score (°)
Diversity metric ⁽¹⁾	Beta(SE)	Р	Beta(SE)	Р
Observed species	0.158 (0.024)	9.55x10 ⁻¹¹	0.086 (0.023)	2.17x10 ⁻⁴
Shannon	0.160 (0.025)	2.16x10 ⁻¹⁰	0.108 (0.023)	3.03x10⁻ ⁶
Simpson	0.082 (0.026)	0.002	0.062 (0.021)	0.003
Chao1	0.060 (0.023)	0.011	0.023 (0.023)	NS
NC not significant. D. 0.00				

NS= not significant: P>0.05.

(1) Standardized to have mean 0, SD 1.

(2) Hippurate associations with diversity metrics adjusted for sex, age, BMI, metabolite batch and family relatedness.

(3) Hippurate diet score associations with diversity adjusted for sex, hippurate, age, BMI, metabolite batch and family relatedness.

6.3.5 OTU and collapsed taxa associations with hippurate

Thirty OTUs and sixteen collapsed taxa (**Figure 6-6**; **Appendix E Tables 1 and 2**) were significantly associated with blood levels of hippurate (*P*<8.61x10⁻⁵ [OTUs]-1.47x10⁻³ [phylum]). At the phylum level, the Firmicutes were associated with reduced levels of hippurate in blood, suggesting reduced diversity. Within the Firmicutes phylum, 28 OTU associations were assigned to the order Clostridiales, though not all associations were in the same direction. Within the order Clostridiales, higher hippurate levels were associated with increased abundances of two OTUs assigned to the family Clostridiaceae and 10 OTUs assigned to the family Ruminococcacea (including 2 *Oscillospira* genus OTUs, 1 *Ruminococcus* genus OTU and 1 *Faecalibacterium prausnitzii* OTU).

Increased abundances of 14 OTUs assigned to the family Lachnospiraceae were associated with reduced hippurate levels. Associations within the Lachnospiraceae family remained strong and in the same direction within the collapsed taxonomies, including at the family level. In particular, within the Lachnospiraceae family there were associations with OTUs assigned to the genera *Ruminococcus* (2 OTUs), *Blautia* (4 OTUs), *Dorea* (2 OTUs) and *Roseburia* (1 OTU), these OTUs were likely driving associations within their respective collapsed genera.

At the class level, lower hippurate levels were associated with increased abundance of Erysipelotrichi, a trend that continued at the order, family and genus levels with

Erysipelotrichales, Erysipelotrichaceae, and *Eubacterium*, respectively. Interestingly there were no OTUs associated to blood hippurate that were assigned to this class.

Outside of the Firmicutes phyla, increased abundances of one OTU assigned to the genus *Actinomyces* was associated with increased blood hippurate levels, this association was likely driving associations within the family Actinomycetaceae and order Actinomycetales collapsed taxa. Increased levels of hippurate were also associated with increased abundances of an OTU belonging to the family Rikenellaceae of the Bacteroidetes phyla and reduced abundances of the collapsed genus *Ralstonia* of the Proteobacteria phyla.

Following a backward stepwise linear regression using P<0.05 as the cut-off threshold, together the remaining OTUs accounted for 58.0% of the variance in Shannon diversity and 7.1% of the variance in hippurate levels (adjusted for diversity).

6.3.6 OTU and taxa associated with both hippurate and the hippurate diet score

Five OTUs and five taxa were associated with hippurate were also associated with the diet score in the same direction (**Table 6-8**). Specifically, reduced abundances of one OTU assigned to the genus *Actinomyces* was associated with increased scores on the diet, similar to hippurate this trend was significant both the family and order levels. At the genus level, reduced abundances of *Ruminococcus* (including 2 OTUs) and *Eubacterium* were associated with increased diet scores were associated to increased abundances of OTUs assigned to the species *Faecalibacterium prausnitzii* and the genus *Clostridiales*. Most associations appeared to be primarily driven by intakes of fruit and whole grains. Though higher abundances of one Clostridiales OTU were associated with increased coffee intake.

I did not identify any significant interactions between the diet score and hippurate/dietassociated microbes independently, nor when abundances were summed, in the prediction of blood hippurate levels.



Figure 6-6. OTU and collapsed taxonomic associations with hippurate Associations between blood hippurate and microbiome variables are represented by the histogram bars on the right side of the plot. The histogram bars represent the $-\log_{10}$ of the *P*-value of the regression and the colour of the bars indicates the direction of association: green, positive; red, negative.

				Hippura	te	Diet sc	ore	
								Foods ⁽³⁾
Order	Family	Genus species	OTU ID ⁽²⁾	Beta (SE)	Р	Beta (SE)	Ρ	<i>P</i> <0.05
Actinomycetales		·	Collapsed	-0.083(0.022)	1.31x10 ^{-₄}	-0.035(0.011)	1.67x10 ⁻³	Fruit: -0.004(0.002) WG: -0.007(0.003)
Actinomycetales	Actinomycetaceae		Collapsed	-0.089(0.021)	2.89x10 ⁻⁵	-0.036(0.011)	1.70x10 ⁻³	Fruit: -0.004(0.002) WG: -0.007(0.003)
Actinomycetales	Actinomycetaceae	Actinomyces	Collapsed	-0.101(0.021)	1.55x10 ⁻⁶	-0.045(0.011)	5.71x10 ⁻⁵	Fruit: -0.005(0.002) WG: -0.008(0.003)
Actinomycetales	Actinomycetaceae	Actinomyces	denovo467	-0.099(0.022)	5.14x10 ⁻⁶	-0.051(0.011)	2.81x10 ⁻⁶	Fruit: -0.005(0.002) WG: -0.009(0.003)
Clostridiales			denovo299	0.113(0.024)	2.21x10 ⁻⁶	0.044(0.010)	9.76x10 ⁻⁶	Coffee: 0.013(0.002)*
Clostridiales	Lachnospiraceae	Ruminococcus	Collapsed	-0.111(0.022)	4.03x10 ⁻⁷	-0.038(0.011)	6.35x10 ⁻⁴	Fruit: -0.005(0.002) WG: -0.008(0.003)
Clostridiales	Lachnospiraceae	Ruminococcus	denovo55	-0.123(0.021)	1.17x10 ⁻⁸	-0.054(0.011)	2.79x10 ⁻⁶	Fruit: -0.006(0.002)* WG: -0.009(0.003)
Clostridiales	Lachnospiraceae	Ruminococcus gnavis	denovo27	-0.107(0.023)	3.04x10 ⁻⁶	-0.064(0.011)	1.99x10 ⁻⁸	Fruit: -0.006(0.002)* WG: -0.009(0.003)
Clostridiales	Ruminococcaceae	Faecalibacterium prausnitzii	denovo469	0.100(0.023)	1.66x10 ⁻⁵	0.034(0.010)	9.24x10 ⁻⁴	WG: 0.007(0.003)
Erysipelotrichales	Erysipelotrichaceae	Eubacterium	Collapsed	-0.083(0.021)	9.30x10 ⁻⁵	-0.040(0.012)	6.12x10 ⁻⁴	Fruit: -0.004(0.002) WG: -0.010(0.003)*

Table 6-8. List of taxa associated with hippurate, the hippurate diet score and foods ⁽¹⁾

*= statistically significant: P<0.0017; WG: whole grain products

- (1) Microbiome OTUs and collapsed taxa significantly associated with both hippurate and the hippurate diet score are shown. Associations were adjusted for covariates (age, Shannon Index, metabolite batch, BMI, sex and family relatedness) and multiple testing using Bonferroni correction. Hippurate diet score associations were also adjusted for hippurate.
- (2) OTU ID assignment is specific to the TwinsUK cohort.
- (3) All foods included in the hippurate diet score were fitted into a backwards stepwise linear regression using P<0.05 as the cut-off threshold with each taxa associated to both hippurate and the diet score. Results displayed are the betas with standard errors of foods at least nominally associated (P<0.05). Statistical significance was defined as P<0.0017 (Bonferronil: 0.05/[10 taxa x 3 foods]).</p>

6.4 Part 2 results

6.4.1 Relationship of diversity, the hippurate trajectory and diet to MetS and its components

Longitudinal hippurate trajectories were significantly associated with Shannon diversity, independently of diet and covariates in a subsample of 1032 twins (15.736[1.96]; P=4.95x10⁻¹⁵), moreover the hippurate trajectory accounted for 6.5% of the variance in Shannon diversity.

Figure 6-7 shows the results of the analysis for associations between diversity, the hippurate trajectory and MetS (**Appendix E Tables 3 and 4** contain the full results). Higher Shannon diversity (R²: 0.0105) and an increasing hippurate trajectory (R²: 0.0054) were associated with a reduced risk of having MetS (**Figure 6-7a**). The variance in MetS attributable to the hippurate trajectory was accounted for 61.1% by Shannon diversity. The variance in MetS attributable to the hippurate trajectory was accounted for 61.1% by Shannon diversity. The variance in MetS 51.9% (Clostridiales OTU) for the five collapsed taxa and 3 of the OTUs that were associated to both hippurate and the diet (**Figure 6-7b**).

Increased Shannon diversity and the hippurate trajectory were significantly associated with a reduced BMI (diversity R^2 : 0.0288) and TG (diversity R^2 : 0.0213; **Figure 6-7a**). Increased Shannon diversity was also associated with higher HDL cholesterol (R^2 : 0.0126). Shannon diversity accounted for 57.6% of the variance in BMI (R^2 : 0.0288) attributable to the hippurate trajectory, the impact of specific OTUs/taxa ranged from 22.6% (*Ruminococcus* genus) to 35.4% (*Faecalibacterium prausnitzii*). Shannon diversity accounted for 63.2% of the variance in TG (R^2 : 0.0087) attributable to the hippurate trajectory, the impact of specific OTUs/taxa ranged from 3.4% (*Eubacterium* genus) to 48.3% (Clostridiales OTU).



hippurate, the diet score and MetS represented as betas with SEs; all variables have been standardized. Figure 6-6b shows the percentage variance the MetS, BMI and triglycerides are attributable to the hippurate trajectory accounted for through each applicable variable.

Abbreviations: MetS, metabolic syndrome; HDL, high density lipoprotein; BMI, body mass index.

6.4.2 Confirmation of results in discordant twins

I identified 55 MZ twin pairs who were discordant (\geq 1 SD apart) for Shannon diversity. Associations between Shannon diversity and hippurate cross-sectionally and the longitudinal trajectory were significant (*P*<0.05), associations with all other variables were in the same direction as in the whole group analysis (**Table 6-9**), except for HDL-cholesterol.

Table 6-9. Associations between hippurate (discovery), the hippurate trajectory, MetS status and components in MZ twins discordant for diversity

Variable	Beta(SE)	Р	R^2
Hippurate (discovery)	0.208 (0.081)	0.013	0.0607
Hippurate trajectory	0.478 (0.078)	9.53x10 ⁻⁸	0.1768
Hippurate diet score	0.149 (0.099)	0.137	0.0136
MetS status*	1.046 (0.299)	0.875	0.0004
BMI	0.021 (0.063)	0.737	0.0010
HDL-cholesterol	-0.035 (0.069)	0.612	0.0022
TG	-0.073 (0.060)	0.233	0.0118

MetS, metabolic syndrome; HDL, high density lipoprotein; TG, triglycerides

A linear regression was conducted using diversity to predict hippurate (discovery), the hippurate trajectory, and MetS status and components in the MZ discordant (1 SD apart in diversity) twin sample.

*Statistical results show the odds ratio. Variables were standardized to have mean=0, SD=1.

6.5 Discussion

In summary circulating levels of hippurate were significantly and positively associated with gut microbiome diversity in all twins and confirmed in MZ twins discordant for diversity. Blood hippurate levels longitudinally and cross-sectionally were also associated with food intakes, specifically self-reported intakes of fruit and whole grains. Longitudinal changes in blood hippurate (reflecting in part food polyphenol content) were also strongly associated with Shannon diversity, accounting for 17.7% of the variance in diversity in MZ discordant twins. Blood hippurate levels and reported intakes of associated foods were associated with 5 OTUs and 5 collapsed taxa independent of diversity. Both an increasing Shannon diversity and higher hippurate trajectory were associated with a lower risk of MetS, which was primarily accounted for by their relationship to BMI and TG, moreover the 5 collapsed taxa and 3 OTUs were associated with MetS.

Metabolite associations with diversity

I identified and validated in an independent sample of male and female twins 5 blood metabolites associated with gut microbiome diversity: hippurate, p-cresol sulfate, phenylacetylglutamine and 3-phenylpropionate which were associated positively; and negatively with hyodeoxycholate. All of these metabolites are derived from or modified by bacterial metabolism.

P-cresol sulfate (also p-cresyl sulfate) and phenylacetylglutamine are potentially toxic uremic solutes formed from the putrefication by colonic bacteria of dietary proteins that have escaped digestion in the small intestine. Our group previously showed p-cresol sulfate and phenylacetylglutamine to be strongly associated with reduced estimated glomerular filtration rate, moreover 52 OTUs assigned to the genera *Christensenellaceae*, *Ruminococcaceae* and *Lachnospiraceae* were associated with phenylacetylglutamine and 3 OTUs assigned to the genera Ruminococcaceae and Lachnospiraceae with P-cresol sulfate (Barrios et al., 2015). Hyodeoxycholate is a secondary bile acid, produced from intestinal bacterial metabolism. In healthy individuals hyodeoxycholate is metabolised by glucuronidation in the human liver and kidneys, a pathway for toxin elimination (Perreault et al., 2013). In individuals with cholestatic liver disease or intestinal malabsorption increased levels of hyodeoxycholate are excreted in the

urine (Sacquet et al., 1983), suggesting hyodeoxycholate may act as a marker of poor liver function. Which is an interesting observation considering the relationship between some metabolic diseases and poor liver function.

Increased blood 3-phenylpropionate and hippurate were associated with increased diversity and also intakes of fruits and whole grains. Both 3-phenylpropionate and hippurate are derived from gut microbial metabolism of polyphenols to benzoates and are significantly correlated (*r*=0.51), though hippurate accounted for a greater degree of variance in diversity therefore I decided to focus on it. Hippurate is emerging as a key mammalian-microbial co-metabolite. It is a glycine conjugate of benzoic acid formed in the mitochondria of the liver (Gatley and Sherratt, 1977) and kidneys (Temellini et al., 1993), through gut bacterial production of benzoic acid from dietary components, primarily polyphenols (Gonthier et al., 2003). Supporting my findings, human feeding studies of foods with a high polyphenol content, such as teas, fruit and coffee, have shown to increase urinary output of hippurate (Gonthier et al., 2003, Walsh et al., 2007). It should be noted however that the ROC analysis showed hippurate was a poor marker of Shannon diversity, therefore future studies should aim to identify biomarkers with better predictive performance.

Microbiome associations with hippurate but not associated with diet

To date, the microbiome profile associated with blood hippurate levels has not been well characterised. Within the TwinsUK dataset I found overall higher hippurate levels were associated to both increased and reduced abundances of 30 OTUs and 16 collapsed taxa, independently of Shannon diversity. At the phylum level, increased levels of hippurate were associated with reduced abundances of Firmicutes. The Firmicutes, which primarily consist of gram-positive bacteria, are one of the dominating phyla of the adult microbiome. One of the first large-scale microbiome studies suggested that a high Firmicutes to Bacteroidetes ratio is characteristic of obesity (Ley et al., 2006), although I did not investigate this ratio. A handful of studies have found reduced Firmicutes levels with increasing BMI (Duncan et al., 2008, Tims et al., 2013, Escobar et al., 2014). Moreover, increased faecal SCFAs have been shown in obese compared to lean subjects which also correlated with a higher ratio of Firmicutes to Bacteroides/Prevotella (Fernandes et al., 2014). Interestingly, polyphenol (concord grape extract) feeding to mice has recently shown to prevent obesity induced by a high fat diet in part by decreasing the Firmicutes/Bacterodetes ratio (Roopchand et al., 2015). Although I did not

find a relationship between total Firmicutes and dietary factors, it is possible our dataset was not powerful enough to detect associations.

Increased hippurate levels were associated to both increased and decreased abundances of bacteria within lower taxonomies of the Firmicutes phyla. Twenty-eight associations with hippurate were with OTUs/taxa belonging to the Clostridiales order which was negatively associated overall. The Clostridiales order belonging to the class Clostridia, are obligate anaerobes found primarily in soil. Directions within lower taxonomic levels of Clostridiales were not consistent with the collapsed order. For instance, increased abundances of two OTUs belonging to the Clostridiales order and 2 OTUs within the Clostridiaceae family were associated with increased blood hippurate levels. OTUs within the Clostridiales order were related to diet and MetS and will be discussed in more detail below.

Notably, the direction of associations between hippurate and OTUs/taxa within the Ruminococcaceae (10 associations) and Lachnospiraceae (14 associations) families of the order Clostridiales, were consistently increased and decreased, respectively. Both Lachnospiraceae and Ruminococcaceae are involved in butyrate production (Vital et al., 2014). In a previous study conducted on the TwinsUK dataset, higher abundances of multiple OTUs assigned to Lachnospiraceae were associated with increased eGFR and reduced levels of blood phenylacetylglutamine (Barrios et al., 2015), a metabolite that was also related to diversity. Another study found that higher abundances of Ruminococcaceae and Lachnospiraceae were associated with increased levels of urinary 3-indoxyl-sulfate and better outcomes in patients at risk of developing gastrointestinal (GI) graft-versus-host-disease (Weber et al., 2015). Moreover, lower abundances of both families are associated with inflammatory bowel disease (Kostic et al., 2014). I did not identify associations with hippurate or other polyphenol-related metabolites in the literature. It is interesting and perhaps noteworthy that I identified clear opposing directions between hippurate levels and these families.

Higher levels of hippurate were associated with reduced abundances of the class Erysipelotrichi, and associations were consistent at the order, family and genus levels with Erysipelotrichales, Erysipelotrichaceae, and *Eubacterium*, respectively. Elevated levels of *Clostridium ramosum*, a member of Erysipelotrichi, have shown in mice fed a high-fat, obesogenic diet, which the authors suspected was due to increasing nutrient absorption (Woting et al., 2014). Moreover, on a 2-month inpatient study of 15 females fed a choline-deficient diet it

was found that faecal levels of of Erysipelotrichi and Gammaproteobacteria were related to changes in liver fat (Spencer et al., 2011).

Outside of the Firmicutes phyla, lower abundances of Actinobacteria were observed with increasing hippurate, these bacteria were further found to be associated with diet and MetS as discussed below. These results provide a microbiome profile associated with hippurate levels although, the complex metabolic relationships require further investigation. As I found diet to be significantly associated with food intakes, I investigated the relationship between these hippurate-associated OTUs and collapsed taxa.

Microbiome associations with hippurate and the hippurate diet

Three of the OTUs and 5 collapsed taxonomies associated to both hippurate and the hippurate diet (independently of diversity), moreover the variance in MetS attributable to the hippurate trajectory that was accounted for by these OTUs and taxa was between 3.7% and 51.9% (**Table 6-10**). These included the increased abundances of the Actinomycetaceae family, *Eubacterium* and *Ruminococcus* being associated with an increased risk (and reduced blood hippurate and diet score) and OTUs within the order Clostridiales and of the species *Faecalibacterium prausnitzii* associated with a reduced MetS risk (and increased blood hippurate and diet score). A detailed discussion of each observation is given below.

Microbiome	Hippurate	Fruit	WG	Coffee	MetS	BMI	TG	HDL
Actinomycetales	$\mathbf{+}$	$\mathbf{\Lambda}$	$\mathbf{+}$		1	1		
Actinomycetaceae	$\mathbf{+}$	$\mathbf{\Lambda}$	$\mathbf{+}$		1	1	1	
Actinomyces	$\mathbf{+}$	$\mathbf{\Lambda}$	$\mathbf{\Lambda}$		↑	1		
Eubacterium	$\mathbf{+}$		$\mathbf{\Psi}$		↑		1	
Ruminococcus	\mathbf{h}	$\mathbf{\Lambda}$	$\mathbf{\Lambda}$		1	1	1	
Ruminococcus (OTU)	\mathbf{h}	$\mathbf{\Lambda}$	$\mathbf{\Lambda}$		1	1	1	1
Clostridiales (OTU)	1			1	$\mathbf{\Lambda}$	$\mathbf{\Psi}$	$\mathbf{\Lambda}$	1
Faecalibacterium prausnitzii (OTU)	^		↑		¥	¥	¥	↑

Table 6-10. Direction of associations between OTUs/collapsed taxa and hippurate, food intakes, MetS and its components.

Abbreviations: WG, whole grain; MetS, metabolic syndrome; BMI, body mass index; TG, triglycerides; HDL, high density lipoprotein cholesterol.

An increasing hippurate trajectory associated with a lower MetS risk

Overall, increased Shannon diversity and the hippurate trajectory were associated with a reduced risk of having MetS. Supporting this, previous studies have shown reduced gut bacterial diversity has been found in metabolic conditions, including atherosclerosis (Koren et

al., 2011), obesity (Turnbaugh et al., 2009) and with phenotypes characteristic of the metabolic syndrome (Le Chatelier et al., 2013). Interestingly, most of the variance in MetS that was attributable to the hippurate trajectory was accounted for by Shannon diversity (61.1%). An increasing hippurate trajectory (accounting for BMI) was similarly associated with a lower BMI and TG (variance 57.6% and 63.2% by Shannon diversity, respectively) at endpoint.

Previous studies that examined urinary excretion or serum levels of hippurate have shown reduced urinary hippuric acid excretion in obesity (Shearer et al., 2008, Waldram et al., 2009, Calvani et al., 2010), though mainly in animal models. To my knowledge, no human studies to date have investigated specifically the relationship between the MetS and markers of dietary polyphenol consumption, such as hippurate. Though the diet score was not associated with MetS, it was associated with a lower BMI for which the hippurate trajectory (60.4%) and Shannon diversity (45.3%) contributed particularly strongly (**Appendix E Table 5**), suggesting that these factors were important mediators between of the effect of diet on obesity and consequential metabolic risk.

Increased abundances of the Actinomycetaceae family, and Eubacterium and Ruminococcus genera associate with MetS risk

The Actinomycetaceae family are typical commensals within the oral cavity. Elevated levels of Actinomycetaceae within the gut have been implicated in conditions where stomach acid production has been compromised, such as with proton pump inhibitor use (Imhann et al., 2016). The metabolic implications of increased abundance of Actinomycetaceae in the gut are not clear at this time, though higher levels within the oral cavity contribute to periodontitis and correlate with reduced insulin sensitivity (Demmer et al., 2015). In rare cases an *Actinomyces* overgrowth contribute to an infection within the gut (abdominal actinomycosis) through forming filamentous branches that grow through damaged mucosal tissue penetrating the gut barrier, forming abscesses and fistula (Bonnefond et al., 2016). As mucosal inflammation and bacterial translocation are emerging as important factors in the aggravation of MetS, *Actinomyces* should be explored further. The relationship between Actinomycetaceae and the foods forming the pupility clear, though these findings suggest a healthy diet may prevent the proliferation of Actinomycetaceae.

Increased abundance of the genus *Eubacterium* was significantly associated with both reduced hippurate and the diet score (in particular whole grain intake), and higher risk of MetS.

The *Eubacterium* genus is within the family Eubacteriaceae of the Firmicutes phyla. Effects at the *Eubacterium* genus level have not been clarified, though different species belonging to the same genus may have different metabolic capacities and therefore be enriched by varying dietary exposures. Contrary to my findings others have shown feeding of whole grains (Martinez et al., 2013) and switching from a Western to plant-based diet (David et al., 2014) have shown to enrich abundances of *Eubacterium rectale*. More studies are needed to outline effects at the *Eubacterium* genus level or with other species.

Higher abundances of the genus Ruminococcus (one OTU and collapsed taxonomy) were associated with MetS risk and in particular higher levels of both HDL and TG. Ruminococcus are most abundant in one of the 3 major, though controversial (Jeffery et al., 2012) enterotypes (Arumugam et al., 2011). Functional analyses have shown the Ruminococcus enterotype to be enriched for genes that produce membrane sugar transporters (Arumugam et al., 2011), which may predispose to weight gain. Ruminococcus are also able to ferment complex carbohydrates (Walker et al., 2011) and produce alcohol (Christopherson et al., 2014). Previous studies have not found the Ruminococcus enterotype to be consistently modified by diet (Wu et al., 2011a, Claesson et al., 2012). I found increased Ruminococcus abundances were shown to be associated with a lower fruit intake. Following 12 weeks feeding of schisandra chinensis fruit, high in flavonoids, to obese women was previously been shown to reduce Ruminococcus abundances (Song et al., 2015). Increased abundances of Ruminococcus were also associated significantly with increased severity of NAFLD lesions (Boursier et al., 2016). Supporting the strong association between TG and Ruminococcus, uncultured phylotypes of the Ruminococcus gnavus-group correlated positively with polyunsaturated serum TGs of dietary origin (Lahti et al., 2013). The Ruminococcus genus contains species with differential effects on metabolism, moreover dietary effects are likely quite complex, for instance increased consumption of foods of animal origin increase abundances of Ruminococcus gnavus, but decrease Ruminococcus bromii and Ruminococcus callidus (David et al., 2014).

Increased abundances of OTUs belonging to the order Clostridiales and Faecalibacterium prausnitzii associate with protection from MetS

Increased abundances of OTUs within the order Clostridiales and species *Faecalibacterium prausnitzii* were associated with hippurate, dietary components and reduced risk of MetS. The

Clostridiales OTU was particularly strongly associated to coffee intake and TG. The gut microbiota play an important role in the metabolism of chlorogenic acids found in high concentrations in coffee. Significantly elevated levels of the *Clostridium coccoides-Eubacterium rectale* group have been shown following the incubation of human faecal microbiota with coffee samples (Mills et al., 2015). Due to the high polyphenol content coffee intake has been thought to have beneficial effects on metabolic health. In a large observational study including 93,179 individuals high coffee intake was associated with a low risk of obesity, MetS and type 2 diabetes, though this was not supported by Mendellian Randomization using five genetic variants associated with coffee intake (Nordestgaard et al., 2015).

Increased abundances of the *Faecalibacterium prausnitzii* OTU were strongly inversely associated with BMI and mildly with higher whole grain intake. *Faecalibacterium prausnitzii* is an important gut commensal accounting for 5% of the total faecal microbiota (Arumugam et al., 2011) that ferments dietary fibre to short-chain fatty acids (including butyrate). In our sample *Faecalibacterium prausnitzii* was strongly associated with Shannon diversity (Beta[SE]: 0.250[0.015]; *P*=5.09x10⁻⁵⁹) and most strongly related to BMI of all microbes related to hippurate and the diet. This clear effect reflects previous findings where *Faecalibacterium prausnitzii* was shown to correlate with microbiome gene count and predict weight loss over time (Le Chatelier et al., 2013). *Faecalibacterium prausnitzii* has previously shown to be depleted in 239 MetS subjects, an effect shown to be partially restored following a long-term (2 years) Mediterranean diet intervention (Haro et al., 2016). Whole grain intake and increased fibre intake as a result allows *Faecalibacterium prausnitzii* to flourish (Benus et al., 2010) and encourages the anti-inflammatory properties of *Faecalibacterium prausnitzii* (Sokol et al., 2008), which may further protect against MetS.

Limitations

There were a number of potential limitations to this study. There were a small number of male subjects in the sample, therefore these results may be applicable only to women. A *de novo* method was used to classify the OTUs into taxonomies. The *de novo* method involves creating a reference panel that is sample specific, therefore replicating our results in independent populations may be difficult to achieve. But this is true of all OTU studies. As the FFQ relies on subject reporting, the accuracy of this data have been called into question. Though in the previous chapters I have shown that reported intakes of those foods significantly associated

with hippurate, coffee, fruit and whole grains, associate with biologically plausible metabolites. If misclassification had occurred it would likely have obscured real findings and not strengthened them. Furthermore, we replicate findings from feeding studies that have shown hippurate urinary excretion to be increased following the consumption of these foods.

There were a number of advantages to this study. I had a large number of unique subjects with metabolomics profiling, dietary information and microbiome profiling. I also had access to a unique longitudinal metabolomics data in order to evaluate the influence of changes in hippurate levels on MetS risk.

Conclusion

In conclusion, hippurate, is associated with gut microbiome diversity and foods high in polyphenols. Subjects whose hippurate trajectory increased showed a reduced risk of MetS and related risk factors, largely accounted for by Shannon diversity. Microbiome OTUs/taxa related to blood hippurate and related food intakes, including increased abundances of Actinomycetaceae, *Eubacterium* and *Ruminococcus* and reduced OTUs within the order Clostridiales and the species *Faecalibacterium prausnitzii* associated with an increased MetS risk. These findings support the gut microbiome as an important mediator in the relationship between diet, in particular high polyphenol foods, and metabolic disease. The potential of hippurate as a marker of alpha-diversity and the interplay between high polyphenol foods, gut microbes and hippurate production should now be established in mechanistic studies ideally under a dietary intervention setting.

Chapter 7 Untangling the Relationship Between Diet and Visceral Fat Mass Through Blood Metabolomics and Gut Microbiome Profiling

In this chapter I created a visceral fat mass diet (VFM) score using the intakes of top foods associated with VFM. I then examined the blood metabolomics and gut microbiome profiles associated with the diet score with the aim of identifying potential biological intermediates linking diet with VFM development.

The work from this chapter has been formatted as a manuscript and submitted to the *International Journal of Obesity*.

7.1 Introduction

Increased visceral fat (VF) is a primary risk factor for cardio-metabolic diseases. Observational studies examining the impact of habitual food consumption on VFM or waist circumference (WC) have found that increased intakes of fruit (Romaguera et al., 2011), dairy products (Romaguera et al., 2011) and related nutrients (Fischer et al., 2015), and whole grains (Caron-Jobin et al., 2011) and fibre (Hairston et al., 2012, Fischer et al., 2015) are associated with lower risk, whereas higher intakes of fried foods and fat (Mollard et al., 2014, Romaguera et al., 2011), alcohol, red and processed meats (Romaguera et al., 2011) and related nutrients (Fischer et al., 2011) and related nutrients (Fischer et al., 2011), and processed meats (Romaguera et al., 2011) and related nutrients (Fischer et al., 2015), sugar-sweetened beverages (Ma et al., 2014, Mollard et al., 2014, Odegaard et al., 2012, Romaguera et al., 2011), refined grains (Caron-Jobin et al., 2011, Romaguera et al., 2011) and high glycaemic index foods (Dal Molin Netto et al., 2014, Romaguera et al., 2010) are associated with higher risk of VF.

Within the past decade, the effects of an unhealthy dietary pattern have been increasingly explored. For instance, Nettleton and colleagues created a diet protective health score using self-reported intakes of whole grains, fish, fruits, vegetables, nuts/seeds (favorable) and red/processed meats, sweets, sugar-sweetened beverages and fried potatoes (unfavorable). They were then able to show that the score was a powerful means to examine gene X diet interactions in obesity in 68,317 subjects of European ancestry (Nettleton et al., 2015). Similarly, Romaguera et al. (Romaguera et al., 2011) formed a summary score

combining the intake frequencies for all foods associated with changed in WC over a 5.5 year median in 48,631 men and women. However, dietary pattern scores have not yet been applied against high-throughput omic datasets.

Metabolomics is being increasingly used in dietary studies in order to further explore the mechanisms of diet on metabolic disease development as extensively shown in this thesis. A recent study from our group showed that some of the metabolites associated with VF mass relate also to type 2 diabetes, insulin resistance and blood pressure (Menni et al., 2016). Moreover, we have found that self-reported reported food intakes are associated with 106 unique metabolites (Pallister et al., 2016), which confirmed the strong role of food intake on metabolic traits (see Chapter 4). At this time, the characteristic metabolomics profile of a metabolically unhealthy diet has yet to be established, nor have those primary metabolites that connect diet to VFM development been defined.

Recent studies are alluding to a role of the gut microbiota in VF development by interacting with food compounds and as a result, supplying metabolites to our bodies (Shoaie et al., 2015), this is supported by in Chapters 4 and 5 where I showed food intakes to be correlated with microbially-derived metabolites. Moreover, I found microbiome alpha diversity and hippurate (a microbial co-metabolite) to be associated with a reduced risk of MetS supporting the interplay of diet, the microbiome and the metabolome in metabolic disease development. These interactions may also intensify the health effects of a calorific, low nutrient dense Western Diet. Through feeding rodents high-fat or high-fat/high-sugar diets, early studies have shown HF feeding increases Firmicutes and reduces Bacteriodetes (Taira et al., 2015) abundances, reduces the abundance of the class Clostridia which was additionally linked to higher visceral adipose tissue (Etxeberria et al., 2015), and increases abundances of sulfidogenic bactera which was associated with higher intestinal inflammation (Shen et al., 2014). Under high fat feeding of animals, cranberry (Anhe et al., 2015) and pomegranate (Neyrinck et al., 2013) polyphenols, resveratrol from red wine (Neyrinck et al., 2013) and glucooligosaccharide (Serino et al., 2012) have been found to be protective of obesity and associated inflammation in by changing the gut microbiome profiles. Also, conjugated linoleic acid, a constituent of dairy fats has been shown to encourage fat loss, which may be due to increasing the abundance of Bacteroidetes (Margues et al., 2015).

Previous studies have not undertaken a multi-omic approach to decipher the influence of an unhealthy diet on VFM, as such the aims of this chapter were:

- To identify foods most strongly associated with VFM from these develop and validate a predictive dietary VFM-risk score.
- ii) To characterise the blood metabolomics profile of the VFM-risk score.
- iii) To characterise the gut microbiome profile of the dietary VFM-risk score.

7.2 Materials and methods

Figure 7-1 shows the summary of the study protocol.

For the analysis I used FFQs completed between 1995 and 2001, in 2007 and 2014 to 2015. I used the 20 food groups as defined in **Section 5.2.1**.

7.2.1 Visceral Fat Mass

During clinical visits a train research nurse or assistant determined visceral fat mass (VFM; g) in 3457 twins using Dual-Energy X-ray Absorptiometry (DXA; Hologic QDR; Hologic, Inc., Waltham, MA, USA) whole-body scanning in the supine position. To analyse the scans, the QDR System Software Version 12.6 was used. VFM was estimated at one cross-section of the whole body (L4-L5), this is the usual area of a computed tomography slice. If VFM was 4 SD above or below the mean VFM, twins were excluded from the analysis. I normalized the VFM data using a rank-based inverse-normalization as it did not follow a normal distribution.

7.2.2 Metabolomic profiling

For this chapter, I used 292 chemically identified metabolites analyzed by the non-targeted Metabolon platform. Quality control of the metabolomics dataset has been described in **Section 3.1.6.1**. For the analysis I included 2218 twins (99.8% female) with metabolomics profiling, BMI and VFM data within \pm 5 years of FFQ completion.

7.2.3 Gut microbiome profiles

16S rRNA gene sequencing was used to determine the faecal bacterial profiles. For a previous study (Goodrich et al., 2014b) microbial DNA was extracted, amplified, sequenced and processed by the Cornell technical team, since this earlier study an additional ~1000 samples were collected and processed using the same protocols. Quality control of the microbiome data were undertaken by Tiphaine Martin, Matthew Jackson and Dr Michelle Beaumont. At 97% sequence similarity, sequencing reads were condensed as operational taxonomic units (OTUs). To do this, UCLUST open-reference clustering was used against Greengenes v13_5 reference within QIIME 1.7.0. Of the total sequences, 6.2% did not cluster to the reference and were therefore excluded from further analysis (Goodrich et al., 2014b).



OTUs that were found in fewer than 25% of individuals were removed from the analysis. Using this threshold and after removing singletons, of 9,840 OTUs only 2,118 OTUs (16%) remained for the final analysis. All OTU counts, including those OTUs present in less than 25% of individuals, were collapsed into taxonomies at the family (124 taxa), genus (283 taxa) and species (153 taxa) levels. For the analysis I only considered taxa fully classified within each taxonomic level. Following rarefaction of the complete OTU table to 10000 reads per sample, alpha-diversity was determined by Shannon's phylogenetic diversity (Faith, 1992) using QIIME. Using linear regression, OTUs were adjusted for technical covariates which included sequencing run and sequence number per sample. I normalized the data using rank-based inverse normalization. I analyzed a subsample (*n*=889) of twins from the FFQ, VFM and metabolomics dataset that additionally had fecal microbiome profiling.

7.2.4 Muther expression data

Gene expression of abdominal fat samples in 825 individuals were analysed with the Illumina Human HT-12 V3 for the Muther study, as described previously (Grundberg et al., 2012). Using random intercept linear regression, 586 individuals were analyzed for AT expression associations with the top metabolite adjusting for age, BMI, metabolite and expression batches, and family relatedness. This analysis was conducted by Dr Cristina Menni.

7.2.5 Statistical analysis

I conducted all statistical analysis using Stata version 12.

7.2.5.1 Food group associations with VFM and diet score formation and heritability

First, I randomly allocated twins to two independent groups: the test (n=1109) and replication (n=1109) groups. I ensured that twin pairs were assigned to the same group. To identify food group associations with VFM, in the test group I ran a linear regression for each of the 20 food groups as predictors of VFM (residual adjusted for BMI), I adjusted for covariates including total fat mass, age, sex, height², family relatedness and DXA batch. The cut-off for statistical significance was defined using Bonferroni correction (P<2.50x10⁻³ = [0.05/20 food groups]). I included food groups that were significantly associated with VFM in the risk score. I calculated the score, first I quartile ranked the consumption frequencies of associated foods and then assigned a score of 0 to 3 by the direction of the association (positive association: Q1=0, Q2=1,

Q3=2, Q4=3; negative association: Q1=3, Q2=2, Q3=1, Q4=0). Next I summed each food group score to create the final risk score that ranged from 0 to 15. I used linear structural equation modelling in Mx (Neale et al., 1992, Neale et al., 2003) to determine the heritability of the VFM diet score (consult **Section 3.2.1** for analysis details).

7.2.5.2 Binary classification test

I next tested the VFM risk score in the replication group by first fitting a logistic regression model with the diet score predicting low and high VFM. I defined the lower tertile of VFM as a negative outcome (0; *n*=369) and the upper tertile of VFM as a positive outcome (1; *n*=370). In the model I adjusted for covariates including total fat mass, age, sex, height², family relatedness, DXA batch, and BMI category (1: <18.5 kg/m²; 2: ≥18.5-24.9 kg/m²; 3: ≥25-29.9 kg/m²; 4: ≥30 kg/m²). To assess the predictive ability of the VFM diet score, I conducted a binary classification test. The ability of the VFM diet score to identify twins with high VFM accurately (sensitivity; true positive rate) and identify twins with low VFM accurately (specificity; true negative rate) was predicted. The receiver operating characteristic curve (ROC) was then generated by plotting the true positive rate against the false positive rate at a number of threshold settings.

7.2.5.3 VFM diet score and metabolomics and microbiome

For each metabolite or microbiome taxon, I ran a random intercept linear regression analysis for each metabolite or microbiome OTU/taxa adjusting for age, BMI, sex and metabolite batch (for metabolomics) or Shannon Index (for microbiome), and family relatedness:

$$Y_{i} = \beta_{0} + \boldsymbol{\beta}_{i} \boldsymbol{X}_{ij} + \delta_{1i} age_{ij} + \delta_{2i} BMI_{ij} + \delta_{3i} Sex_{ij} + \gamma_{1i} Z_{ij} + \zeta_{j} + \varepsilon_{ij}$$

where Y_i is the metabolite/microbiome OTU/taxa, X_{ij} is the VFM diet score of twin *j* from pair *i*, Z_{ij} is the metabolite batch or Shannon Index and ζ_j , is the family-specific error component that captures the unobserved heterogeneity or family characteristics. I also used Bonferroni correction to account for multiple testing giving a significance threshold for metabolites of 1.71×10^{-4} (0.05/(292 known metabolites)). For the microbiome analysis I defined statistical significance within each taxonomic level, **Table 7-1** shows the thresholds by level.

A large number of potentially correlated metabolites were associated with the VFM diet, therefore I next undertook a backwards stepwise linear regression including all associated metabolites, using a cut-off threshold of P<0.01.

	VFM diet score		VFM	
Level ⁽¹⁾	Number of variables	Р	Number of variables ⁽²⁾	Р
Family	124	4.03x10 ⁻⁴	2	0.025
Genus	283	1.77x10 ⁻⁴	2	0.025
Species	153	3.27x10 ⁻⁴	2	0.025
OTU	2118	2.36x10 ⁻⁵	8	6.25x10 ⁻³
	(A) D. (1 90 9	

Table 7-1. Statistical significance thresholds for microbiome analysis using Bonferroni correction

(1) Bonferroni correction was calculated within each level.

(2) Variables significantly associated with the VFM diet score were tested for their association against VFM.

7.2.5.4 VFM and metabolomics and microbiome

To determine if the remaining metabolites, and OTUs/taxa (both residual-adjusted for BMI) associated with the VFM diet score were also associated with VFM, I used each of these variables as predictors of VFM (residual-adjusted for BMI at scan) in a linear regression including the covariates scan age, sex, total fat mass, height², scan batch and metabolite batch (for metabolomics) or Shannon Index (for microbiome), family relatedness and for the VFM diet score.

$$VFM_{i} = \beta_{0} + \beta_{1i}Yi + \beta_{2i}X_{ij} + \delta_{1i}age_{ij} + \delta_{3i}Sex_{ij} + \delta_{4i}Total_fat_mass_{ij} + \delta_{5i}Height_squared_{ij} + \delta_{6i}Scan_batch_{ij} + \gamma_{1i}Z_{ij} + \zeta_{j} + \varepsilon_{ij}$$

where Y_i is the metabolite/taxon, X_{ij} is the VFM diet score of twin *j* from pair *i*, Z_{ij} is the metabolite batch for metabolomics data or Shannon Index for microbiome data and ζ_j , is the family-specific error component that captures the unobserved heterogeneity or family characteristics. Associations which passed the Bonferroni cut-off were considered significant. For metabolites the threshold was 5.56×10^{-3} (0.05/9 metabolites). The assignment for statistical significance for the microbiome analysis is shown in **Table 7-1**.

I identified MZ twins who were discordant (1 SD apart) for VFM, this included 80 pairs in the metabolomics dataset and 27 pairs in the microbiome subsample. In this group, I ran a linear regression to validate top associations between diet, metabolites and the microbiome and VFM.

7.2.5.5 Top microbiome and metabolite associations with food groups

To confirm the associations between the metabolites/microbiome and the VFM diet score were not due to intakes of other foods, I ran a backward stepwise linear regression model including the VFM diet score and food groups not forming the score to predict the associated metabolite and OTU/taxa. I used a cut-off threshold of P<0.01 for metabolites and P<0.05 for microbiome.
I then assessed whether the foods contributing to the VFM diet score were independently driving the association between the VFM diet score and taxa/metabolites. I fitted all 20 food groups into a backward stepwise linear regression model using cut-off thresholds of P<0.01 for metabolites and P<0.05 for microbiome.

7.2.5.6 VFM diet score association with VFM mediated by metabolite and microbiome

I first determined the proportion of the variance of VFM attributable to the VFM diet score after accounting for all covariates (age, sex, BMI, total fat, height², family relatedness, metabolite batch, Shannon Index and scan batch). This quantity was represented by r_x^2 . I next determined the proportion of the variance in VFM explained by the VFM diet score by adjusted for the same covariates as above but additionally including the metabolite or OTU/taxon (r_{xy}^2). To calculate the percentage of the VFM diet score association that was mediated by the metabolite or OTU/taxa (r_y^2), I determined the proportion of the variance of VFM that is due to the VFM diet score association with the metabolite or OTU/taxa: $1 - (r_{xy}^2/r_x^2)$.

7.2.5.7 Microbiome association with VFM mediated by metabolite

I determined the proportion of the variance of VFM attributable to the taxon after taking into account all covariates as above in addition to the VFM diet score (r_x^2) . I next determined the proportion of the variance in VFM explained by the taxon after adjusting for the same covariates as above but additionally adjusting for the metabolite (r_{xy}^2) . To calculate the percentage of the taxon association that was mediated by the metabolite (r_y^2) , I determined the proportion of the variance of VFM that is due to the metabolite association with the taxon: $1 - (r_{xy}^2/r_x^2)$.

7.3 Results

7.3.1 VFM food group associations

The characteristics of the sample as well as the training and validation groups are shown in

Table 7-2.

Table 7-2. Study population characteristics

	Whole (<i>n</i> =2218)	Training (<i>n</i> =1109)	Validation (<i>n</i> =1109)
	Mean (SD)	Mean (SD)	Mean (SD)
Age (years)	58.3 (10.9)	58.1 (11.1)	58.4 (10.8)
BMI (kg/m²)	26.2 (4.8)	26.1 (4.8)	26.4 (4.8)
Total fat (grams)	6827 (2180)	6779 (2178)	6911 (2258)
Visceral fat (grams)	562.7 (293.7)	553.9 (293.8)	572.7 (295.5)
Height (m)	1.61 (0.06)	1.62 (0.06)	1.62 (0.06)
Sex (M:F)	4:2214	0:1109	4:1105
Food intakes (servings/week)			
Vegetables	34.3 (15.4)	33.8 (15.0)	34.8 (15.8)
Fruit and fruit juices	22.4 (13.3)	22.7 (13.8)	22.1 (12.8)
Nuts and legumes	7.9 (5.5)	7.8 (5.6)	8.0 (5.4)
Whole grains	10.2 (7.9)	10.2 (7.7)	10.2 (8.1)
Refined grains	8.4 (7.4)	8.4 (7.4)	8.5 (7.4)
Fermented dairy	6.3 (4.8)	6.4 (4.7)	6.3 (4.8)
White meat	1.9 (1.3)	1.9 (1.3)	2.0 (1.3)
Seafood	2.4 (2.1)	2.4 (1.9)	2.5 (2.2)
Red, processed meat and eggs	6.9 (4.0)	6.9 (3.9)	6.9 (4.0)
Fried and fast foods	4.5 (3.4)	4.5 (3.4)	4.6 (3.4)
Sweets and sweet baked products	15.7 (13.6)	16.2 (13.3)	15.2 (13.8)
Chocolate	4.0 (5.8)	4.1 (6.0)	3.9 (5.7)
Butter and cream	4.4 (6.6)	4.7 (6.8)	4.1 (6.4)
Spreads and dressings	8.3 (8.4)	8.4 (8.8)	8.2 (8.1)
Soy and other milks	0.2 (0.8)	0.2 (0.8)	0.2 (0.8)
Milk	3.9 (4.2)	3.8 (4.1)	4.0 (4.3)
Soda	2.0 (5.2)	1.9 (5.5)	2.1 (4.9)
Coffee	8.7 (10.4)	8.8 (10.5)	8.6 (10.4)
Теа	19.3 (13.7)	19.5 (13.8)	19.1 (13.6)
Alcohol	6.0 (8.0)	5.7 (7.8)	6.3 (8.3)

No significant differences between the training and validation sets.

I found 5 food groups were significantly associated with VFM in the training dataset (**Table 7-3**). Higher intakes of fruit, whole grain and fermented dairy products were associated with reduced VFM, whereas increased intakes of fried and fast foods and red, processed meat and eggs were associated with increased VFM.

Food group	Bota	SE	D
	Dela	5L	·
Fruits	-0.005	0.001	1.95x10 ^{-₅}
Red, processed meat and eggs	0.016	0.005	3.94x10 ⁻⁴
Fermented dairy products	-0.011	0.004	1.14x10 ⁻³
Fried and fast foods	0.015	0.005	1.18x10 ⁻³
Whole grains	-0.008	0.002	1.27x10 ⁻³
VFM diet score ⁽²⁾	0.033	0.006	7.38x10 ⁻⁹

Table 7-3. List of food groups significantly associated with VFM and the VFM diet score association with VFM in the training group $^{(1)}$

 20 food groups were tested for their association with VFM in the training group adjusting for covariates (scan batch, age, BMI, total fat, height², sex and family relatedness) and multiple testing (*P*<0.0025).

(2) The VFM diet score was formed from quartile ranking the VFM-associated food groups, scoring them according to the direction of association and summing the scores.

Based on the direction of the food group association with VFM in the training dataset, I created the VFM diet score and tested it in the validation sample. I found 93.21% of the subjects were accurately classified into low and high VFM by the diet score. Moreover, the sensitivity (true positive rate) and specificity (true negative rate) of the VFM diet score were 93.72% and 92.70%, respectively. The ROC curve is displayed in **Figure 7-2**. The area under the receiver operating curve (AUC) was 0.9841 (95% CI: 0.9772; 0.9911). Notably, I found the association between the diet score and VFM to be significant in the VFM-discordant MZ twin sample (0.281[0.091]; *P*=0.002).



The best-fitting model for the heritability analysis of the VFM diet score was the AE model, with a heritability estimate of 44% (95% CI: 37%, 50%) **(Table 7-4** shows details of heritability analysis).

				log-			
Model	Α	С	E	likelihood	X ²	Р	AIC
	0.44	0.00	0.56				
ACE	(0.31, 0.50)	(0.00, 0.11)	(0.50, 0.63)	9067			5493
	0.44		0.56				
AE*	(0.37, 0.50)	-	(0.50, 0.63)	9067	0.00	Incalc	5491
		0.33	0.67				
CE	-	(0.27, 0.38)	(0.62, 0.73)	9090	23.40	0.000	5514
			1.00				
E	-	-	(1.00, 1.00)	9192	124.70	0.000	5613

 Table 7-4. Results of the structural equation modelling for heritability of the visceral fat

 mass diet score

*Best fitting model; Abbreviations: A, additive; C, common; E, environmental; df, degrees of freedom; AIC, Akaike's information criterion; Incalc, incalculable.

Structural equation modeling was performed on the visceral fat mass diet score. The best-fitting model is indicated by the lowest AIC.

Figure 7-3 displays the nutrient profile of the VFM diet score (**consult Appendix F Table 1 for details**). Individuals scoring highly on the VFM diet score had significantly increased intakes of fat (notably monounsaturated fatty acids) and reduced carbohydrate (notably, sucrose and non-starch polysaccharides [NSP]), vitamin C and magnesium intakes.



7.3.2 VFM diet score metabolomics associations

Thirty metabolites were significantly associated ($P < 1.71 \times 10^{-4}$) with the VFM diet, shown in **Table 7-5**. Overall, these metabolites belonged to 21 unique sub-pathways, specifically, there were associations with 12 lipids, 5 amino acids, 5 vitamins and cofactors, 3 carbohydrates, 3 xenobiotics, 1 nucleotide and 1 peptide.

After I adjusted for other food group intakes (**Table 7-5**) all associations between the VFM diet score and metabolites remained (*P*<0.01) however 6 associations did not pass adjustment for multiple testing (primarily lysolipids).

7.3.3 Metabolites associated with the VFM diet score and food groups independently

Following the backward stepwise regression including all food groups, I found that 18 metabolites were significantly (P<3.33x10⁻⁴ (0.05/[5 food groups x 30 metabolites])) associated with the food groups forming the VFM diet score (**Table 7-5**). Fruit intakes were associated with 11 metabolites, suggesting it was an important influence on associations between the VFM diet score and metabolites. Moreover, 5 metabolites were significantly associated with whole grain intake, 3 metabolites with red, processed meat, and eggs, and 2 metabolites with fried and fast food intakes. I found associations between the food groups forming the VFM diet score and the 12 remaining metabolites did not pass statistical significance.

Table 7-5. List of metabolites significantly associated with the VFM diet score and with each food group independently (P<0.01 backware)	ſd
stepwise regression)	

				VFM Score adjusted			
		VFM SC	ore	toods		Foods associated V	
Metabolite	Sub-pathway	Beta (SE)	Ρ	Beta (SE)	Ρ	<i>P</i> <0.01	<i>P</i> <3.33x10 ^{⁻⁴}
Eicosapentaenoate	Essential fatty acid	-0.058(0.007)	3.24x10 ⁻¹⁷	-0.052(0.006)	2.22x10 ⁻¹⁵	Fruit (0.005(0.002)) FD (0.011(0.004)) FF (-0.017(0.006))	WG (0.011(0.003))
Indolepropionate	Tryptophan metabolism	-0.056(0.007)	1.58x10 ⁻¹⁶	-0.054(0.007)	2.91x10 ⁻¹⁶	Fruit (0.006(0.002)) WG (0.008(0.003)) RM (-0.018(0.005))	
3-Carboxy-4-methyl-5-propyl-2- furanpropanoate	Fatty acid, dicarboxylate	-0.055(0.007)	2.43x10 ⁻¹⁶	-0.045(0.006)	3.19x10 ⁻¹²	FF (-0.023(0.007))	WG (0.013(0.003))
Docosahexaenoate	Essential fatty acid	-0.054(0.007)	6.17x10 ⁻¹⁶	-0.045(0.006)	1.30x10 ⁻¹²	WG (0.010(0.003))	Fruit (0.008(0.002))
Stachydrine	Food component, Plant	-0.056(0.007)	7.95x10 ⁻¹⁶	-0.057(0.007)	2.59x10 ⁻¹⁶	RM (-0.019(0.006))	Fruit (0.014(0.002))
3-Phenylpropionate	Phenylalanine & tyrosine metabolism	-0.059(0.007)	1.70x10 ⁻¹⁵	-0.053(0.008)	3.53x10 ⁻¹²	Fruit (0.006(0.002))	WG (0.014(0.003)) FF (-0.028(0.007))
Hippurate	Benzoate metabolism	-0.052(0.007)	1.84x10 ⁻¹³	-0.052(0.007)	5.06x10 ⁻¹⁴		Fruit (0.010(0.002)) WG (0.011(0.002))
Catechol sulfate	Benzoate metabolism	-0.051(0.007)	1.75x10 ⁻¹²	-0.046(0.007)	1.94x10 ⁻¹⁰	WG (0.008(0.003))	Fruit (0.008(0.002))
Glycerate	Glycolysis, gluconeogenesis, pyruvate metabolism	-0.049(0.007)	3.81x10 ⁻¹²	-0.042(0.007)	1.57x10 ⁻⁹		Fruit (0.010(0.002))
Pyridoxate	Vitamin B6 metabolism	-0.049(0.007)	9.08x10 ⁻¹²	-0.042(0.007)	2.96x10 ⁻⁹		Fruit (0.007(0.002)) WG (0.011(0.002))
Threitol	Nucleotide sugars, pentose metabolism	-0.044(0.007)	7.73x10 ⁻¹¹	-0.043(0.007)	1.65x10 ⁻¹⁰		Fruit (0.011(0.002))
Butyrylcarnitine	Fatty acid metabolism (also BCAA metabolism)	0.040(0.007)	4.77x10 ⁻⁹	0.039(0.007)	9.09x10 ⁻⁹		RM (0.020(0.005))
alpha-Hydroxyisovalerate	Valine, leucine and isoleucine metabolism	0.037(0.007)	1.12x10 ⁻⁷	0.027(0.006)	4.48x10 ⁻⁵		RM (0.022(0.005))
1-Arachidonoylglycerophospho- ethanolamine*	Lysolipid	0.038(0.007)	2.09x10 ⁻⁷	0.030(0.007)	3.43x10 ⁻⁵	WG (-0.011(0.003)) FD (-0.013(0.005))	
Threonate	Ascorbate and aldarate metabolism	-0.036(0.007)	4.51x10 ⁻⁷	-0.033(0.007)	3.66x10 ⁻⁶		Fruit (0.007(0.002))
1,5-Anhydroglucitol (1,5-AG)	Glycolysis, gluconeogenesis, pyruvate metabolism	0.035(0.007)	1.80x10 ⁻⁶	0.026(0.007)	3.46x10 ⁻⁴		Fruit (-0.008(0.002))
Uridine	Pyrimidine metabolism, uracil containing	-0.032(0.007)	2.05x10 ⁻⁶	-0.03(0.007)	9.21x10 ⁻⁶	WG (0.008(0.003))	
Stearidonate	Long chain fatty acid	-0.033(0.007)	2.65x10 ⁻⁶	-0.028(0.007)	3.20x10 ⁻⁵	None	
1- Docosahexaenoylglycerophosphocholin e*	Lysolipid	-0.032(0.007)	2.95x10 ⁻⁶	-0.027(0.007)	4.14x10 ⁻⁵	FD (0.014(0.004))	

Table 7-5. List of metabolites significantly associated with the VFM diet score and with each food group independently (*P*<0.01 backward stepwise regression)

		VFM Sc	VFM Score adjusted VFM Score ⁽¹⁾ foods ⁽²⁾			Foods associated ⁽³⁾		
Metabolite	Sub-pathway	Beta (SE)	Ρ	Beta (SE)	Ρ	<i>P</i> <0.01	<i>P</i> <3.33x10 ⁻⁴	
Bilirubin (Z,Z)	Hemoglobin and porphyrin metabolism	-0.033(0.007)	3.88x10 ⁻⁶	-0.032(0.007)	4.69x10 ⁻⁶		FF (-0.020(0.006))	
1-Oleoylglycerophosphoethanolamine	Lysolipid	0.032(0.007)	9.21x10 ⁻⁶	0.023(0.007)	1.35x10 ⁻³	WG (-0.008(0.003))		
1-Arachidonoylglycerophosphocholine*	Lysolipid	0.032(0.007)	1.76x10⁻⁵	0.024(0.007)	1.26x10 ⁻³	WG (-0.010(0.003))		
Pantothenate	Pantothenate and CoA metabolism	-0.029(0.007)	4.20x10 ⁻⁵	-0.029(0.007)	6.28x10 ⁻⁵	None		
4-Androsten-3beta,17beta-diol disulfate 1*	Sterol, Steroid	0.032(0.008)	4.67x10 ⁻⁵	0.019(0.007)	7.86x10 ⁻³	None		
X-11793oxidized bilirubin*	Hemoglobin and porphyrin metabolism	0.028(0.007)	5.10x10 ⁻⁵	0.027(0.007)	7.01x10 ⁻⁵	WG (-0.009(0.002)) FD (-0.015(0.005))		
trans-4-Hydroxyproline	Urea cycle; arginine-, proline-, metabolism	0.027(0.007)	8.28x10 ⁻⁵	0.031(0.007)	7.13x10 ⁻⁶	FD (-0.013(0.005))	RM (0.028(0.006))	
Glycoursodeoxycholate	Bile acid metabolism	0.032(0.008)	1.00x10 ⁻⁴	0.028(0.008)	5.70x10 ⁻⁴	None		
gamma-Glutamylvaline	gamma-glutamyl	0.023(0.006)	1.11x10 ⁻⁴	0.023(0.006)	1.38x10 ⁻⁴	FD (-0.014(0.004))	Fruit (-0.005(0.001))	
1-Eicosatrienoylglycerophosphocholine*	Lysolipid	0.028(0.007)	1.32x10 ⁻⁴	0.020(0.007)	4.40x10 ⁻³	WG (-0.009(0.003))		
Proline	Urea cycle; arginine-, proline-, metabolism	0.028(0.007)	1.44x10 ⁻⁴	0.028(0.007)	1.45x10 ⁻⁴	FF (0.015(0.006))	Fruit (-0.008(0.002))	

FD: Fermented dairy; FF: Fried and fast foods; RM: Red meat; WG: Whole grain products

(1) Metabolite associations with the VFM diet score were adjusted for covariates (batch effects, age, BMI and sex) and multiple testing (P<1.71x10⁻⁴).

(2) The VFM diet score and 15 food groups not forming the score were fitted into a backward stepwise linear regression model to predict each significant metabolite using *P*<0.01 as the cut off threshold.

(3) All 20 food groups were fitted into a backward stepwise linear regression model to predict each significant metabolite using P<0.01 as the cut off threshold. Significant results shown only for foods forming the VFM diet score. Associations passing the Bonferonni cut-off were considered statistically significant: P<3.33x10⁻⁴ = 0.05/[5 food groups x 30 metabolites].

7.3.4 Metabolites associated to both the VFM diet and VFM

I included all 30 metabolites in a backward stepwise linear regression and found 9 metabolites were significantly and independently associated with the VFM diet score (**Table 7-6**), these metabolites accounted for 14% of the variance in the VFM diet score. Notably, four of these metabolites also passed the cut-off four multiple testing (P<5.56x10⁻³). I also found 6 out of the 9 metabolites were nominally associated (P<0.05) with both the VFM diet and VFM. The remaining 3 metabolites were associated only with the VFM diet and not VFM.

I found higher scores on the VFM diet score and high VFM (independently of diet and total body fat) were associated with lower hippurate and bilirubin (Z,Z), and higher alphahydroxyisovalerate and butyrylcarnitine (**Table 7-6**). Notably, in the VFM-discordant MZ twins, associations between VFM and butyrylcarnitine (0.199[0.087]; *P*=0.023) and hippurate (-0.297[0.095]; *P*=0.002) were significant (**Figure 7-4**; **Appendix G Table 2**). On average, these four metabolites accounted for 18.5% (range: 13.5% [alpha-hydroxyisovalerate]-28.9% [hippurate]) of the variance in the association between the VFM diet score and VFM (**Table 7-6**). Table 7-6. List of metabolites independently associated with the VFM diet score (P<0.01 in backward linear regression), their association with VFM and the proportion of the association of the VFM diet score with VFM that is mediated by the VFM diet score association with the metabolites (P<5.56x10⁻³)

	VFM diet score stepwise ⁽¹⁾ VFM ⁽²⁾					% association	
Metabolite name	beta(SE)	Р	beta(SE)	Р	diet R ² no metabolite ⁽³⁾	diet R ² with metabolite ⁽⁴⁾	through metabolite
Hippurate	-0.45(0.10)	2.15x10 ⁻⁵	-0.081(0.012)	1.33x10 ⁻¹¹	0.0312	0.0222	28.8%
alpha-Hydroxyisovalerate	0.38(0.10)	9.60x10 ⁻⁵	0.050(0.013)	1.65x10 ⁻⁴		0.0270	13.5%
Butyrylcarnitine	0.33(0.10)	8.54x10 ⁻⁴	0.072(0.013)	5.86x10 ⁻⁸		0.0267	14.4%
Bilirubin (Z,Z)	-0.31(0.10)	1.76x10 ⁻³	-0.049(0.013)	1.88x10 ⁻⁴		0.0258	17.3%
Indolepropionate	-0.33(0.11)	2.21x10 ⁻³	-0.030(0.012)	1.40x10 ⁻²			
1-Arachidonoylglycerophospho- choline*	0.27(0.10)	5.20x10 ⁻³	0.031(0.012)	1.07x10 ⁻²			
Eicosapentaenoate (EPA; 20:5n3)	-0.75(0.10)	1.13x10 ⁻¹³	0.020(0.012)	NS			
Threonate	-0.32(0.11)	2.59x10 ⁻³	-0.016(0.012)	NS			
X-11793Oxidized bilirubin*	0.33(0.11)	2.63x10 ⁻³	-0.004(0.012)	NS			

NS= not significant: P>0.05

(1) Thirty metabolites significantly associated with the VFM diet score (Table 7-5) were adjusted for covariates (batch effects, age, BMI and sex) and fitted into a backward stepwise linear regression to predict the VFM diet score using *P*<0.01 as the threshold cut-off.

(2) Nine metabolites independently associated with the VFM diet score were tested for their association with VFM adjusted for covariates (age, batch effects, BMI, total fat, sex, height², and family relatedness). Associations passing the Bonferonni cut-off were considered significant (*P*<5.56x10⁻³).

(3) the proportion of the variance in VFM explained by the VFM diet score after taking into account all covariates (age, sex, BMI, height², and batch effects).

(4) the proportion of the variance in VFM explained by the VFM diet score after taking into account all covariates as in (1) and adjusting for the metabolite.



7.3.5 VFM diet score microbiome associations

In a subsample of 889 twins I found higher scores on the VFM diet were associated with lower gut microbiome diversity (Shannon Index; -0.025[0.009], P=6.26x10⁻³), after I adjusted for VFM this association was reduced but remained significant (-0.020[0.010], P=0.035).

I found the VFM diet score was associated significantly with 8 OTUs (**Table 7-7**) and 6 taxa (**Table 7-8**). However, after I adjusted for intakes of other food groups, the associations became nominally significant (P<0.05) (**Table 7-8**).

7.3.6 Microbiome taxa associated to both the VFM diet and VFM

I found higher abundances of the species species *Eubacterium dolichum* (0.057[0.019], $P=2.73\times10^{-3}$) was associated significantly with increased VFM and a *Bifidobacterium* OTU (OTU ID: 4426298; -0.046[0.016], P=0.005) with reduced VFM (adjusting for the VFM diet score). I found that 16.4% of the effect of the VFM diet score on VFM ($r_x^2=0.0238$) was mediated by *E. dolichum* ($r_{xv}^2=0.0199$) and 17.2% by the *Bifidobacterium* OTU.

7.3.7 Eubacterium dolichum and hippurate associated with both VFM and VFM diet

I tested associations between those 4 metabolites significantly associated to both VFM and the VFM diet to determine if *E. dolichum* and the *Bifidobacterium* OTU may be related to the metabolomics profile. I found that higher abundances of *E. dolichum* were associated with lower levels of hippurate at the nominal level (*P*<0.05) after adjusting for VFM, the VFM diet score, Shannon Index and covariates (-0.075[0.032], *P*=0.021). Moreover, I found that 36.9% of the effect of *Eubacterium dolichum* on VFM (r_x^2 = 0.0065) was mediated by hippurate (r_{xy}^2 = 0.0041) independently of diet and covariates.

7.3.8 Hippurate association with adipose tissue transcriptome

Adipose tissue gene expression levels in the greater twin population, in order to provide potential mechanisms of the effect of hippurate on fat mass development. Higher levels of hippurate were associated with increased expression of neuroglobin (0.016[0.004], $P=9.82 \times 10^{-6}$). Neuroglobin has roles in cellular energy maintenance as a vertebrate globin family member.

Table 7-7. List of OTUs associated with the VFM diet score (unadjusted and adjusted for other food intakes), their association with foods forming the VFM diet score and their independent association with the VFM diet score (*P*<0.05 in backward linear regression)

		VFM score ⁽¹⁾		VFM score ac	ljusted foods ⁽²⁾	Foods associated ⁽³⁾
OTU ID	Assigned Taxonomy	Beta(SE)	Р	Beta(SE)	Р	<i>P</i> <0.05
4426298 ⁽⁴⁾	k_Bacteria; p_Actinobacteria; c_Actinobacteria; o_Bifidobacteriales; f_Bifidobacteriaceae; g_Bifidobacterium; s_	-0.058(0.011)	6.19x10 ⁻⁷	-0.053(0.011)	9.56x10 ⁻⁷	FD (0.029(0.008))* FF (-0.022(0.010))
183686	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_; s_	0.053(0.011)	1.25x10 ⁻⁶	0.042(0.011)	9.01x10 ⁻⁵	RM (0.018(0.009)) WG (-0.016(0.004))*
592616	<pre>k_Bacteria; p_Firmicutes; c_Erysipelotrichi; o_Erysipelotrichales; f_Erysipelotrichaceae; g_; s_</pre>	0.050(0.011)	3.35x10 ⁻⁶	0.042(0.010)	4.48x10 ⁻⁵	RM (0.027(0.009))*
2368865	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_; g_; s_	-0.047(0.011)	1.11x10 ⁻⁵	-0.044(0.010)	2.40x10 ⁻⁵	FD (0.031(0.006))*
509709	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; g_; s_	-0.051(0.011)	1.15x10 ⁻⁵	-0.047(0.011)	2.57x10 ⁻⁵	RM (0.011(0.004)) WG (-0.023(0.008))
New.0.Reference OTU51	k_Bacteria; p_Bacteroidetes; c_Bacteroidia; o_Bacteroidales; f_Rikenellaceae; g_; s_	0.049(0.011)	1.44x10 ⁻⁵	0.045(0.011)	3.87x10 ⁻⁵	WG (-0.011(0.004))
3801267	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Veillonellaceae; g_Veillonella; s_parvula	-0.043(0.011)	1.75x10 ⁻⁵	-0.039(0.010)	7.29x10 ⁻⁵	Fruit (0.006(0.002))
2407149	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; g_; s_	-0.043(0.010)	2.24x10 ⁻⁵	-0.028(0.010)	0.006	None

*= statistically significant: *P*<0.0025; FD: fermented dairy; FF: fried and fast foods; RM: red meat; WG: whole grain products

(1) OTU associations with the VFM diet score were adjusted for covariates (age, Shannon Index, BMI and sex) and multiple testing (*P*<2.36x10⁻⁵ [Bonferroni: 0.05/2118 OTUs]).

(2) The VFM diet score and 15 food groups not forming the score were fitted into a backward stepwise linear regression model to predict each significant OTU using *P*<0.05 as the cut off threshold.

(3) All 20 food groups were fitted into a backward stepwise linear regression model to predict each significant OTU using *P*<0.05 as the cut off threshold. Significant results shown only for foods forming the VFM diet score.

(4) OTU 4426298 is the only taxon associated with VFM independently of the VFM diet score (Beta[SE]: (-0.046[0.016]; P=0.005).

Table 7-8. List of taxa associated with the VFM diet score (unadjusted and adjusted for other food intakes), their association with foods forming the VFM diet score and their independent association with the VFM diet score (*P*<0.05 in backward linear regression)

		VFM Score ⁽¹⁾		VFM Score	Foods associated ⁽³⁾	
Taxon	Level	beta(SE)	Р	beta(SE)	Р	<i>P</i> <0.05
Actinomyces	genus	0.052(0.011)	9.77x10 ⁻⁷	0.052(0.011)	9.77x10 ⁻⁷	FF (0.028(0.009))* RM (0.027(0.008))*
Lachnospira	genus	-0.045(0.009)	2.79x10 ⁻⁶	-0.038(0.010)	8.33x10 ⁻⁵	Fruit (0.006(0.002))
Actinomycetaceae	family	0.043(0.011)	5.47x10 ⁻⁵	0.043(0.011)	5.47x10 ⁻⁵	FF (0.021(0.010)) RM (0.024(0.008))
Eubacterium dolichum ⁽⁴⁾	species	0.042(0.011)	8.47x10⁻⁵	0.043(0.011)	6.19x10 ⁻⁵	WG (-0.010(0.004))
Veillonella dispar	species	-0.039(0.011)	3.05x10 ⁻⁴	-0.031(0.011)	4.00x10 ⁻³	None
Anaeroplasmataceae	family	-0.037(0.010)	3.75x10 ⁻⁴	-0.036(0.010)	3.37x10 ⁻⁴	Fruit (0.007(0.003)) WG (0.011(0.004))

*= statistically significant: P<0.0025: FF: Fried and fast foods; RM: Red meat; WG: Whole grain products

(1) Taxa associations with the VFM diet score were adjusted for covariates (age, Shannon Index, BMI and sex) and multiple testing.

(2) The VFM diet score and 15 food groups not forming the score were fitted into a backward stepwise linear regression model to predict each significant taxon using *P*<0.05 as the cut off threshold.

(3) All 20 food groups were fitted into a backward stepwise linear regression model to predict each significant taxon using *P*<0.05 as the cut off threshold. Significant results shown only for foods forming the VFM diet score.

(4) Eubacterium dolichum is the only taxon associated with VFM independently of the VFM diet score (Beta[SE]: 0.057[0.019], P=2.74x10⁻³).

7.4 Discussion

In this chapter, I developed a dietary VFM risk score and evaluated it against blood metabolomics and faecal microbiome profiles. I characterised the metabolomics profile of the VFM dietary pattern and additionally found a specific gut bacterial species to associate with this pattern and VFM independently of confounders such as age, BMI and total fat mass. In particular, using this novel dataset I found the species *E. dolichum* in the gut and hippurate in blood may connect an unhealthy diet to VFM.

The score I created was strongly predictive of VFM in the TwinsUK cohort, and it was replicated VFM-discordant MZ twins. This allowed me to further explore the potential effects of diet on VFM development through metabolomics and microbiome methods. Overall, I found 4 metabolites were significantly associated with both the VFM diet score and VFM (after adjusting for diet). These metabolites also accounted for some portion of the variance in the association between the VFM diet score and VFM. Specifically lower levels of hippurate and bilirubin (Z.Z), and higher levels of alpha-hydroxyisovalerate and butyrylcarnitine were associated with higher VFM diet scores and VFM. Both alpha-hydroxyisovalerate and butyrylcarnitine are metabolites involved in BCAA catabolism and fatty acid metabolism, both have previously been shown to be higher in obese children (Butte et al., 2015) and adults (Moore et al., 2014). In particular, alphahydroxyisovalerate has been found to predict insulin resistance and glucose intolerance well (Gall et al., 2010, Varvel et al., 2014). Higher levels of both alpha-hydroxyisovalerate and butyrylcarnitine were associated with increased intakes of red and processed meats and eggs. Although animal sources of fats and protein have yet to be connected to deranged BCAA metabolism in humans, a previous study has shown through feeding mice a HF diet the addition of BCAA (which are primarily sourced from animal-derived proteins) aggravates insulin resistance by activating the mTOR kinase pathway (Newgard et al., 2009).

Bilirubin is an endogenous anti-oxidant that also has roles in haemoglobin and porphyrin metabolism. Similar to my findings, reduced levels of serum bilirubin have been associated with increased abdominal adiposity and metabolic derangements (Wu et al., 2011b, Kwon et al., 2011, Jenko-Praznikar et al., 2013). I found increased reported consumption of fried and fast foods were associated with reduced bilirubin (Z,Z). A previous study identified reported intakes of total fatty acids were correlated with reduced bilirubin in serum (Jenko-Praznikar et al., 2013). This association may be potentially derived from elevated levels of

oxidative stress depleting bilirubin. In support of this, increased intakes of fried and fast foods were associated with reduced levels of EPA and increased proline (related to collagen breakdown (Palka, 1996)) that might suggest higher inflammation and oxidative stress (Krishnan et al., 2008). Moreover, vegetable oil frying also lowers polyphenols and was shown to increase liver microsomal lipid peroxides when fed to mice (Quiles et al., 2002).

The primary metabolite connecting diet to VFM was hippurate. Overall, reduced levels of hippurate in blood were associated with higher VFM, a finding that I additionally replicated in VFM-discordant MZ twins, the VFM dietary score and E. dolichum in the gut. Hippurate is a glycine conjugate of benzoic acid formed in the mitochondria of the liver (Gatley and Sherratt, 1977) and kidneys (Temellini et al., 1993), but is also a is a mammalian-microbial co-metabolite formed by gut bacterial metabolism of dietary components, in particular polyphenols (Gonthier et al., 2003, Walsh et al., 2007), to produce benzoic acid. I found increased levels of hippurate to be independently associated with higher intakes of fruit and whole grain products, which are high in polyphenols. Concerning obesity and metabolic diseases animal models which have shown reduced hippuric acid urinary excretion in obesity (Shearer et al., 2008, Waldram et al., 2009, Calvani et al., 2010) and higher levels in Type II diabetes (Williams et al., 2005) compared to controls. To my knowledge no human studies to date have extensively evaluated the relationship between VFM and blood metabolite biomarkers of dietary polyphenol consumption, including hippurate. Although in the previous chapter I found hippurate to be positively associated with microbiome diversity as well as inversely associated with MetS risk. Interestingly, I also identified higher levels of hippurate in blood to be associated with increased expression of neuroglobin in adipose tissue. Neuroglobin, is a globin protein expressed mainly in neurons and some endocrine tissues (Burmester et al., 2000), it has roles in protecting cells against hypoxia and oxidative stress (Burmester et al., 2007). Hypoxia, which is a lack of oxygen reaching tissues, has emerged in recent studies as an important mechanism in the progression of adipose tissue dysfunction (Kim et al., 2014). This finding suggests a mechanism in which hippurate could protect against adipose tissue dysfunction and resultant VFM development.

E. dolichum is a bacterial species belonging to the family *Erysipelotrichaceae*, increased abundances of *E. dolichum* were associated with higher dietary VFM scores and VFM (independently of diet). This finding suggests this microbe may contribute to VFM development through diet, notably lower intakes of whole grains. Similar to my findings, in a

murine model of Western-style diet induced obesity, abundances of E. dolichum were increased (Turnbaugh et al., 2008), additionally further metagenomics analysis showed the E. dolichum genome may be enriched for genes that allow this microbe to have an advantage under a Western-style diet as it is enriched for phosphotransferase proteins which have roles in the import and metabolism of simple sugars. An additional study of Japanese quail strains (atherosclerotic-resistant and non-resistant) showed increased E. dolichum abundance in atherosclerotic-resistant quails compared to control when fed a high cholesterol diet (Liu et al., 2015). I believe this is the first human study to find a connection between E. dolichum and a diet high in fat and low in fibre and micronutrients however I do not believe there to be literature on the mechanisms of the effect of this species on VFM development. It may be that the relationship between E. dolichum and VFM is primarily an artefact of poor diet and not specifically contributing to VFM, however the association was unaffected when adjusted for the diet score. Within TwinsUK higher abundances of E. dolichum have associated with increased scores on the frailty index, a finding that remained strong after adjustment for diet and diversity (Jackson et al., 2016b). The association I found between E. dolichum and hippurate is difficult to untangle with my limited dataset, though should be assessed in future studies.

This chapter had many strengths. I believe this to be the first large-scale study applying multi-omic methods to explore the effect of diet on VFM. The food intake associations with VFM replicate findings from previous studies, this confirms the strength and validity of the dietary VFM score. In particular, I replicated associations from previous studies finding higher reported intakes of fruit (Romaguera et al., 2011), dairy products (Romaguera et al., 2011), and whole grains (Caron-Jobin et al., 2011) with reduced VFM or WC, while higher intakes of fried and fatty foods (Mollard et al., 2014, Romaguera et al., 2011) and red and processed meats (Romaguera et al., 2011) with increased VFM or WC. Similar to VFM (Direk et al., 2013), I found the dietary risk score determined strongly by genetics (h^2 : 44%) this is in line with previous studies suggesting the heritability of an 'unhealthy' diet pattern ranges from 33 to 50 % (Pallister et al., 2014). I also experienced a number of limitations throughout this chapter. The study population was mainly females and my results may not apply to men. Due to the cross-sectional nature of the study, I cannot attribute cause and effect to my findings. I adjusted for a large number of potential confounders, though I still face possible residual or unmeasured confounding due to other unmeasured factors or measurement error. Although, as I did adjust for a large set of confounders as well as for multiple testing it is not likely that these factors

would totally account for my results. The measurement of the gut microbiome was further limited by using 16S gene sequencing. Subsequent investigations using more accurate metagenomic methods may allow for a more thorough examination into microbe-metabolite interactions at the functional level. I used a variety of time points for sampling, however it is likely my results may be enhanced if the same time point was used throughout. I also did not have repeated measurements for individuals and therefore I could not evaluate intra-individual variation. Finally, I was not able to replicate my findings in an independent population, despite this I was able to replicate top association in MZ VFM-discordant twins that are matched for age, gender and the baseline genetic sequence.

Conclusions

An unhealthy dietary is an important influence on VFM. In this chapter I connected a dietary VFM score and VFM to a single gut microbial species and blood metabolites. In particular, I found the species *E. dolichum* seems to link a dietary pattern with low in fruit, whole grains and fermented dairy product intakes and high red, processed meat and eggs, and fried and fast foods intakes to VFM, which was independent of BMI and total fat mass. Additionally, I found hippurate, an important benzoate and microbial co-metabolite, to link these features to the microbiome. Finally, I found hippurate to associate with neuroglobin expression in adipose tissue, alluding to a potential mechanism of interaction. In the future, studies may aim to verify my findings through dietary intervention and establish further how our findings influence long-term metabolism and health.

Metabolomics has emerged from this work as a key application for furthering nutrition research. Through examining the metabolome in a bodily system, specific organ, tissue or cell, information is provided as to the metabolic state of an individual. Nutrition and the metabolome are highly related as nutrients and non-nutrient food constituents supply chemical components to the body, although separating these contributions can be challenging (Gibney et al., 2005). In spite of this, using self-reported food intakes candidate biomarkers have been identified within the metabolome in this study and others (O'Sullivan et al., 2011, Altmaier et al., 2011, Guertin et al., 2014b), validating this approach. However, the genetic effect on the metabolite profile is highly variable (Suhre et al., 2011), as such application of the co-twin control method can provide an excelled way to further metabolomics studies by providing cases and controls matched for age, sex and genetics.

In this thesis, through using metabolomics in collaboration with self-reported dietary intakes I have identified a number of candidate biomarkers that can be used and assessed in subsequent epidemiology studies. I evaluated the utility of using metabolite scores to assess food intakes and applied the scores to study the odds of developing MetS. I used metabolites to investigate the metabolomics signature of gut microbiome diversity and connected increased levels of hippurate in blood to intakes of foods high in polyphenols, diversity and lower odds of developing MetS. Finally, I attempted to untangle the influence of a diet predictive of VFM on VFM development and found blood levels of hippurate and *E. dolichum* in the gut may be important links. Throughout each chapter I used discordant MZ twins who are controlled for age, sex and the baseline genetic sequence to validate my findings. In the discussion below I have highlighted the key findings from each chapter.

Key points/learning from the thesis

Chapter 4: Metabolite-food intake discovery analysis

- Self reported food intakes correlate strongly with levels of blood metabolites.
- Of 601 total metabolites assessed, 128 metabolites (21.3%) were associated with food intakes and dietary patterns using stringent cut-offs for multiple testing.
- Wine intake and seafood/fish intakes yielded the strongest and most abundant results, many replicated from previous studies.

- 72 food intake-metabolite associations were novel: including associations between mushroom intake and ergothioneine, apples/pear and 3-phenylpropionate, and meat intake and trans-4-hydroxyproline.
- Assessing both dietary patterns and specific food group intakes yielded unique results.

Chapter 5: Metabolite-food scores

- Creating metabolite scores provided an advantage over using single metabolites to measure food intakes.
- Differences between each of the methods to create the scores were negligible, though the weighted method performed the best.
- Metabolite scores performed moderately for alcohol (AUC>0.8) and (AUC>0.7) for fruit, whole grains, seafood, fried foods, tea and coffee.
- Metabolite scores for seafood, whole grains, and butter and creams were associated with a reduced odds of future MetS, a finding not replicated by intake data.

Chapter 6: Metabolite markers of gut microbiome diversity

- Gut microbiome alpha-diversity was associated with five blood metabolites: hippurate, p-cresol sulfate, phenylacetylglutamine and 3-phenylpropionate which were associated positively; and negatively with hyodeoxycholate
- Hippurate, a benzoate metabolite derived from polyphenols, was associated with intakes of fruit and whole grains cross-sectionally and longitudinally.
- Hippurate was positively associated with OTUs and collapsed taxa within the family Ruminococcacea.
- Hippurate was negatively associated with OTUs and collapsed taxa within the Firmicutes phyla, the family Lachnospiraceae, the class Erysipelotrichi, and the order Actinomycetales.
- Longitudinal hippurate levels were associated with a reduced odds of MetS, which was largely mediated by Shannon diversity (61.1%).

Chapter 7: Dietary VFM score and metabolomic and microbiome profiles

- Creating a diet score was a robust means to predict VFM
- The dietary risk score had a strong metabolomics signature: associating with 30 metabolites
- Notably, in blood lower hippurate and bilirubin (Z,Z) levels, and higher alphahydroxyisovalerate and butyrlcarnitine levels were associated to increased VFM and the VFM diet score.
- Increased abundances of the collapsed taxonomy assigned to *Eubacterium dolichum* was associated with both increased VFM and the VFM diet score
- Increased *Eubacterium dolichum* adbundances were associated with reduced blood hippurate and adipose tissue expression of neuroglobin, a protein protective of hypoxia and oxidative stress.

Chapter 4: Metabolite-food intake discovery analysis

I explored all metabolomic associations with reported food intakes and dietary patterns and used MZ twins discordant for food intakes to validate the findings. Overall, I identified and validated 128 metabolites to be associated between self-reported food intakes and diet patterns. To my knowledge, 72 metabolite associations with food intakes were novel (have not been identified in blood previously by studies also using FFQ data (Guertin et al., 2014b, Zheng

et al., 2014)). Identifying this many novel associations is encouraging for future dietary metabolomics studies suggesting there is much potential within the metabolome. Moreover, that I replicated many findings from previous studies further supports the methods used. Interestingly, 21 metabolites were associated to dietary patterns not items which may suggest that through measuring dietary patterns we are able to capture information not obtained by measuring single foods independently. In my analysis, I did not attempt to assess the utility of using single food items versus combined biomarkers of dietary patterns when applied to other indicators of health such as MetS, fasting blood glucose, waist circumference or BMI, which could be an area for future studies. Certainly a larger number of metabolites representing diverse metabolic pathways are associated with dietary patterns highlighting this potential area of importance. Although, my results with principal components patterns are population-specific and therefore cannot be replicated, which is why I chose to analyse the Mediterranean diet using an a priori method. This method yielded a lower number of associations than the comparable Fruit and Vegetable PC pattern, though this also highlights that the Mediterranean diet score has limitations, which have been outlined previously (Hoffman and Gerber, 2013). Moreover, I did not find any metabolite-associated SNPs to be related to food intake, though more precise food intake data or using combined genetic risk may have a stronger effect. Moreover, testing these SNPs or combined genetic risk SNPs to examine gene-diet interactions are a potential area for future study. I used discordant MZ twins as a validation population which was very stringent criteria, though this strongly supports the value of the associations I identified. Future large-scale feeding studies need to be undertaken to confirm our associations, to properly quantify effects and also to determine the origin of the associations (e.g. whether the metabolite is derived from a food itself and its quantity in food). Further adding to the complexity of associations are intermediate processes such as one's genetic makeup and the influence of gut bacteria metabolism. Finally, 32 metabolites that were associated to food intakes and 29 to dietary patterns are currently not identified but in the future may provide useful metabolic insights.

Chapter 5: Metabolite-food scores

For the Chapter 5 analysis, I created metabolite scores (when there were multiple metabolites associated to a food) using three methods (quartile ranking, continuous scoring and weighted) and compared each method against the top associated metabolite. I believe this was the first

time this was done before and certainly with such a large dataset. In the training dataset, I was able to form metabolite scores for fifteen different food groups (vegetables, fruit, whole grains, nuts and legumes, seafood, white meat, red meat, fried foods, sweets and sweet baked products, butter and cream, spreads and dressings, milk, tea, coffee and alcohol). To form these groups I used highly stringent methods: associations must have passed multiple testing cut-offs, been replicated in MZ discordant twins and maintained significance following the data reduction steps. Of course, the methods I used may have been too stringent and I may have formed more powerful groups if I had used more metabolites, on the other hand, if my results were to be replicated this would be easier done with fewer metabolites. Nonetheless, I replicated my results in the testing group, therefore at least some of these scores should translate as important markers in independent populations. As for the performance of the scores, they generally performed better than the top single metabolite and the weighted method tended to perform best although differences were negligible. At predicting high and low consumers in the test group, metabolite scores performed well (AUC>0.8) for alcohol and reasonably (AUC>0.7) for fruit, whole grains, seafood, fried foods, tea and coffee. Moreover, I applied a unique method by using food group liking as an alternative way of indicating food intakes, I found metabolite scores for all groups but vegetables, white meat and spreads and dressings to correlate with reported food liking which also supported the results of the ROC analysis. Based on my findings from this analysis, each scoring method appears to be suitable for use in biomarker studies, though due to its simplicity, the continuous method (multiplying metabolite concentrations according to the association direction and summing) may be ideal, especially in studies using multiple populations.

In the second part of the chapter, I used validated metabolite scores to examine their association with MetS and its components in the twins at least 5 years after metabolomics profiling. I found high levels of metabolites correlated to consumption of seafood, whole grains and butter and cream intakes to associate with a lower risk of MetS. Although reported intakes did not reflect these findings in our sample. However, the dataset I used to examine these associations was very small and only 25 twins developed MetS in later years. Therefore, further analysis could be done on dataset with a larger number of cases. Although, the ability to identify an association with MetS with such a small sample size might be encouraging for the usage of metabolomics for future studies. Also, the potential applicability for these metabolite biomarkers

to be used as pre-disease markers of dietary risk in a personalised medicine setting is an exciting area for further development.

Chapter 6: Metabolite markers of gut microbiome diversity

I found five blood metabolites correlated with gut microbiome diversity (hippurate, p-cresol sulfate, phenylacetylglutamine and 3-phenylpropionate which were associated positively; and negatively with hyodeoxycholate) and examined their association with diet and relationship with MetS. Figure 8-1 shows a summary of the findings. Overall, higher levels of hippurate in blood were the strongest correlate of increased gut microbiome diversity, which was validated in MZ twins discordant for diversity. Hippurate was also associated with reported food intakes and I therefore examined it more thoroughly. Hippurate is a metabolite derived from benzoic acid which is primarily produced by gut bacterial metabolism of dietary phenolic acid (Gonthier et al., 2003). Supported by previous studies (Gonthier et al., 2003, Walsh et al., 2007), reported intakes of high polyphenol-containing foods, whole grains, fruits and coffee, were significantly predictive of longitudinal changes in blood hippurate levels. In the faecal microbiome I characterised a gut microbiome profile correlated with blood hippurate levels and overall identified 46 associations. This included higher hippurate levels being associated with lower levels of Firmicutes phyla overall, and lower and higher levels of Lachnospiraceae and Ruminococcaceae, respectively. Interestingly, both Lachnospiraceae and Ruminococcaceae are butyrate-producing bacteria (Vital et al., 2014), therefore measuring faecal short-chain fatty acid production may be an interesting area future investigation. Comparing those OTUs/taxa associated with hippurate, I found 5 OTUs and 5 collapsed taxa to be associated to the the hippurate diet score as well, suggesting food intake may modify hippurate production through enriching these bacteria, though this would need to be confirmed by feeding studies. I had the advantage of using longitudinal hippurate data and found an increasing trajectory to be associated with a reduced odds of later having MetS, reduced BMI and TG. The longitudinal metabolite data certainly provided a strong advantage to this analysis, though Shannon diversity accounted for 61.1% of the variance in the association between the hippurate trajectory and MetS, suggesting that Shannon diversity was strongly driving this association. Although, the dietary factors were not associated with MetS, which may be related to less measurement precision. Untangling the dietary influence of hippurate and Shannon diversity on MetS and its components is challenging due to the nature of FFQ data. It is likely that food intakes are

influencing diversity, the gut microbiome profile and hippurate production though separating the effect of diet is a challenge. It is important to consider whether there are diet-microbe interactions in the formation of hippurate, I attempted to investigate this but did not find any associations, potentially due to the limited precision of the dataset but also the time between sample collection and FFQ completion may be of relevance. It is likely some individuals have a gut microbiome profile with an increased capacity for hippurate production and therefore increased ability to handle oxidative stress, a central component of metabolic diseases. However no clear patterns of microbes emerged. Future dietary intervention studies should test our findings by feeding individuals high polyphenol-containing foods and testing the changes induced on the microbiome, and blood hippurate levels. Longer-term studies should explore the influence of hippurate on MetS.



Figure 8-1. Summary of findings from Chapater 6

Figure 8-1a represents increased gut microbiome diversity correlates with increased blood hippurate. Figure 8-1b represents increased intakes of high polyphenol foods (whole grains, coffee and fruits) predict an increasing hippurate trajectory. Moreover, an increasing hippurate trajectory correlates with lower risk of MetS (metabolic syndrome) at endpoint. Images have been obtained and modified from google search under the usage right option 'labelled for reuse with modification'.

Chapter 7: Dietary VFM score and metabolomic and microbiome profiles

I identified foods most strongly associated with VFM and developed and validated a predictive dietary VFM-risk score. I also characterised the blood metabolomics and gut microbiome profiles associated with the dietary VFM-risk score and VFM. **Figure 8-2** shows a summary of the findings. I found reported intakes of fried and fast foods and red, processed meats and eggs

were associated with increased VFM while intakes of fruits, fermented dairy products, and whole grain products were associated with lower VFM, moreover the dietary risk score was highly predictive of VFM correctly identifying 92.7% of twins in the upper and lower tertiles of VFM. The dietary risk score had a strong metabolomics signature, associating with 30 metabolites, many of which were associated to food intakes in the first and second chapters. After data reduction, I found 4 metabolites were associated with both increased VFM and the VFM-diet, including reduced hippurate and bilirubin (*Z*,*Z*) (involved in hemoglobin and porphyrin metabolism), and increased alpha-hydroxyisovalerate (involved in valine, leucine and isoleucine metabolism) and butyrylcarnitine (involved in fatty acid and BCAA metabolism). It is encouraging that hippurate was found to associate with future MetS in Chapter 6 and also with VFM, as waist circumference is a component of MetS.

The VFM diet score was also associated with 6 collapsed taxa and 8 OTUs, within the faecal microbiome, though only 1 Bifidobacterium OTU and 1 taxa, Eubacterium dolichum was associated with both VFM and the VFM-diet. Interestingly, increased levels of E. dolichum associated with a reduced level of hippurate in blood. Upon further investigation I found increased hippurate in blood to be associated with elevated adipose tissue expression of neuroglobin (Burmester et al., 2000) a protein which protects cells against hypoxia and oxidative stress (Burmester et al., 2007), the importance of hypoxia in the development adipose tissue dysfunction has recently emerged (Kim et al., 2014). This may also suggest that hippurate protects against adipose tissue dysfunction and VFM development as a result. It would be useful if mechanistic studies were performed to confirm if hippurate plays a protective role in adipose tissue dysfunction and also, importantly, determine the function of neuroglobin in adipose tissue as no studies have outlined this before. Abundance of the species Eubacterium dolichum has previously shown to increase in mice fed a Western-style diet high in saturated fat and simple carbohydrates and low in plant polysaccharides (Turnbaugh et al., 2008). This study on mice found E. dolichum to function in the import and processing of simple sugars (Turnbaugh et al., 2008), using metagenomics analysis of the microbiome data from my study could be used to confirm their findings (Turnbaugh et al., 2008). To my knowledge this is the first time a relationship between Eubacterium dolichum and a poor diet has been made in humans and the usefulness of this marker for epidemiology and clinical studies when diet histories are difficult to obtain should be evaluated.



Figure 8-2. Summary of findings from Chapter 7

Figure 8-2 represents the visceral fat mass (VFM) diet which is composed of high intakes of fried and fast foods and red processed meat and eggs and low intakes of fruits, whole grains and fermented dairy products. The VFM diet and VFM correlate with lower levels of blood hipurate and faecal *Eubacterium dolichum*. The color of the arrows represent direction protective of VFM: *red*, increased VFM; *green*, lower VFM. Images have been obtained and modified from google search under the usage right option 'labelled for reuse with modification'.

What I learned and challenges

In general I found using FFQ data throughout the thesis to be problematic. Though I did have success in finding associations, to bring the field forward we will need to step away from using FFQs and begin to apply intervention studies or use more detailed dietary data (such as food records). I also found defining food intake groups to be problematic. Although it is a common practice in nutritional epidemiology, statistically intakes of the foods included within these food groups are not always correlated and therefore I found providing a logical reasoning for group assignment or understanding the biological mechanisms for the metabolite associations was challenging. Moreover, the relative importance of a single food as opposed to dietary patterns is difficult to ascertain. I did learn how to conduct a principal component dietary pattern analysis, though whether these patterns are more reflective of patterns in reporting rather than true behaviours is a question that requires asking. I recognise that FFQ data is the best data I had access to for the study and for some foods it is better at capturing habitual consumption for (e.g. coffee and alcohol).

I also learned that there are many ways to analyse metabolomics data. In the field of metabolomics it is a common practice to perform dimension reduction techniques (such as PCA or partial least squares regression) due to the correlation between metabolites. The results of my thesis may have been quite different if I had first undertaken dimension reduction techniques. For one, I would have had less risk for false negative associations with fewer variables. However, it is likely that not all of the metabolites included in latent variables (e.g. PCs) would be directly related to the intake of a food, similar to the issue that not all foods included in a food group are likely to explain a metabolite association. Had I undertaken dimension reduction it would also have been challenging to unravel the reasoning for correlations between certain metabolites.

Throughout my work I learned and employed the co-twin control method that was a unique aspect of the thesis as a whole. One issue I did have was with defining discordance for food intake data. As FFQ data is not precise using a 1 SD cut-off for discordance may not have been an ideal definition for all foods. The significant food-metabolite associations I found in discordant MZ twins may have just been a reflection of improved accuracy of the FFQ to measure certain food intakes. It may have been better to define MZ discordance as one twin consuming a food while the other does not, though likely there would have been fewer instances of these and it would have been food-dependent. The twin model does have potential for further exploitation especially in the context of dietary intervention studies, where twin studies are few. Despite all of these challenges I encountered, I do believe that metabolomics has strong potential for improving our understanding of the relationship between diet and disease and we should continue the work being done in this field.

Future directions and conclusion

Dietary metabolomics research is still very much in its infancy and there are a number of issues which need to be overcome before the ideal clinical test panel of dietary biomarkers to measure food intake status can be achieved. To take the field forward, I have specific recommendations regarding the work I undertook in this thesis (providing Chapter 7 as an example) and general recommendations concerning the field as a whole.

With regards to carrying the work I undertook in this thesis forward, I would recommend the following:

- 1. Replication of significant results in independent cohorts using similar datasets
- 2. Repeating the analysis with new metabolomics platforms
- 3. Dietary intervention:
 - a. To confirm and quantify the food intake effect on biomarkers

b. To evaluate metabolomics and microbiome findings with the metabolic phenotypes studied (VFM and MetS [plus components])

Due to the very good predictive performance of the VFM dietary risk score, I would suggest undertaking a 4- or 5-week dietary intervention modelled after the score. Ideally, this could also be done as a cross-over study where for one phase a VFM-risk diet is consumed while during the other phase a VFM-protective diet is consumed, separated by a washout period. Therefore, for the VMF-risk phase, the diet would consist of low intakes of fruit, whole grains and fermented dairy products and increased intakes of fried and fast foods and red, processed meat and eggs, and the opposite for the VFM-protective phase. A challenge may be determining the ideal quantities of foods to be consumed and the incorporation of other foods to ensure a balanced diet. This could involve collecting subject food records prior to beginning the study and personalising the diets to ensure compliance and real-world applicability of the findings, though this would utilize more time and resources. Visceral fat mass, faecal microbiome and blood metabolomics data would be measured at the beginning and end of each phase.

To analyse the data I would aim to confirm the findings from my analysis in the greater twin dataset. Therefore I hypothesize that following a diet high in fried and fast foods and red, processed met and eggs, and low in fruit, whole grains and fermented dairy products will increase VFM, alpha-hydroxyisoverate, butyrlcarnitine and *E. dolichum*, and reduce levels of hippurate and bilirubin (Z,Z). However, if the same technologies were used it could be interesting to confirm if overall results using all metabolites and microbiome phenotypes are comparable to my findings in the larger twin dataset. Moreover, twin modelling could be an asset, for instance to determine if MZ twins are more alike than DZ twins in their response to the intervention, or through using the co-twin control method, which in this case could be confirming the hypothesis using MZ twins discordant for feeding (e.g. comparing endpoint phenotypes of one twin fed the VFM-risk diet compared to baseline phenotypes of other twin).

General areas to further nutritional metabolomics studies

For the future of nutritional metabolomics there are a number of issues to be resolved, some of which I encountered, and areas of research which require further exploration to strengthen the field. For instance, many of the associations I identified were with metabolites endogenous to the human body and therefore determining the origin of these metabolites and quantifying the

effect of food consumption on levels is complex. The food metabolome potentially contains many metabolites that are exogenous to the human body and therefore are less influenced by metabolic processes, though it may be more difficult to obtain biological information on these metabolites. Delving deeper into the food metabolome therefore may be useful, this would also involve compiling more complete metabolomics profiles of foods into databases, which is now already taking place (Scalbert et al., 2014), such as the FooDB (www.foodb.ca), Phenol-Explorer (www.phenol-explorer.eu) and PhytoHub (www.phytohub.eu). Moreover, careful intervention studies are needed to evaluate the utility of biomarkers, whether they are applicable for short (hours/days) or long-term usage (months/years), as well as the relevance of the type of biosample provided (e.g. urine vs blood). Feeding studies evaluating short-term biomarkers within exposure are of course the easiest to undertake whereas longer-term studies will be more challenging due to difficulties such as participant compliance and cost factors, however these will be more valuable.

The generalisability of my findings to both males and females and determining racial effects have also not been thoroughly explored here nor in many other studies. One study found one third of metabolites were influenced by sex, with network analysis revealing sex-specific metabolic pathways (Krumsiek et al., 2015). Racial effects have not been well characterised, though one study found distinct differences between the metabolomics profiles of Caucasians versus African Americans in response to atenolol treatment (Wikoff et al., 2013), I did however confirm some of the top findings from two food metabolomics studies on African Americans (Zheng et al., 2014).

Longitudinal data of metabolite levels, which I was able to have access to for hippurate, will likely provide stronger correlations with intakes and power for outcome prediction, improving studies more still. In additions, repeating the findings of my study with more advanced platforms measuring more metabolites and with increased quantitative accuracy (such as NMR-based metabolomics) will likely generate further novel findings. Applications of food intake biomarkers to study disease outcomes is one of the primary aims of the research and future studies should begin to apply these methods, for which longitudinal data will be a particular asset.

Interactions between the human gut microbiome and diet and the subsequent influence on metabolism are not well characterised at this time therefore future studies are needed. One way this could be taken further is through re-analysing the gut microbiome data used in my study using metagenomics data to determine the metabolic functions of the bacteria associated

with hippurate or the VFM diet. Analysing the faecal metabolome would add further information. Another continuing issue with microbiome data is that methods have not been effectively established, for instance studies are quite heterogeneous with regards to quality control procedures and referencing panels used making cross-study comparisons difficult. Even within my thesis, two different referencing methods were used for chapters 6 and 7. This meant that it was difficult to compare results across the thesis, it may have been useful if I had repeated the analysis using the same datasets for both chapters. To aid this and other issues collaboration between cohorts is essential.

Conclusion

In conclusion, I have shown that metabolomics has great promise for furthering nutrition research. Using high-throughput omic data and a unique human cohort I have generated a number of novel and exciting findings. In particular, I found unique correlations between food intakes and metabolites, I evaluated different methods for creating combined metabolite scores, I identified hippurate as a useful blood marker of gut microbiome diversity that is also related to diet and MetS and VFM. I also characterised the metabolomics and microbiome profile of an unhealthy diet associated with VFM. I have shown the broad applications of metabolomics to nutritional epidemiology which should stimulate further work in this area.

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Appendix A. Chapter 1 Appendices

Table 1. Sel	ection of twin	studies es	timating heritab	ility of energy a	nd macro	onutrient int	takes (froi	n Palliste	r et al., 20	14).*		
								Heri	itability			
			Dietary		Twin			Ma	acronutrie	nt intake	•	
			assessment		pair	Energy	Gr	ams per da	ay	P	ercent ene	rgy
Author	Sex	Age	method	Twin registry	count	intake	Fat	Protein	СНО	Fat	Protein	СНО
Pimpin <i>et al.</i> (2013) ^a	Male + female (<i>n</i> = 3605)	21 months	3-day diet diary	Gemini cohort (UK)	384 MZ 832 DZ	12 ^b	11	12	9	10	8	9
Liu <i>et al.</i> (2013)	Male + female (<i>n</i> = 358)	11-13 (x̄ = 11.8)	3-day diet diary	USC Twin Study (US)	94 MZ 85 DZ	48	44	31	43			
Faith <i>et al.</i> (1999) ^⁵	Male + female (<i>n</i> = 108)	≥ 18	Two Buffet-style meals	New York Obesity Centre twin registry (US)	36 MZ 18 DZ	33						
Wade <i>et al.</i> (1981) ^c	Female (<i>n</i> = 46)	19-58 (x̄= 39.2)	3-day diet diary	Toronto Twin Register (Canada)	13 MZ 10 DZ	11 <i>n</i> s	_f	20 <i>ns</i>	66	48 ns	70	67
Heller <i>et al.</i> (1988) ^c	Male + female $(n = 400)$	17-66 (x = 36)	4-day food diary	Australia	106 MZ 94 DZ	38 <i>n</i> s	24 <i>ns</i>	8 <i>ns</i>	31 <i>ns</i>			
	· · · ·	、 ,					SFA: 10 <i>ns</i>		SC: 20 <i>ns</i>			
							MUFA: 33 <i>ns</i>		CC: 55			
							PUFA: 3 <i>n</i> s					
de Castro <i>et</i> <i>al.</i> (1993) ^d	Male + female (<i>n</i> = 390)	x = 38.8	7-day diet diary	Minnesota Twin Registry, (US)	109 MZ 86 DZ	65 (65)	51 (47)	57 (58)	61 (64)			
Hasselbalch et al. (2008)	Male & female (<i>n</i> = 1212)	18-67 (x = 38)	247-item FFQ	Geminakar (Denmark)	600							
	Males Females					38 32				36 ⁹ 53 ⁹	28 55 ⁹	36 49
Hur <i>et al.</i> (1998) ^e	Male + female (<i>n</i> = 335)	18-77 (x = 42.4)	67-item FFQ	MSTRA (Multi- national)	66 MZ 51 DZ	32 (40) ^e	35	16 <i>ns</i>	25			
							SFA: 37		SC: 24 <i>ns</i>			
							PUFA: 46		CC: 18 <i>ns</i>			

FFQ, food frequency questionnaire; USC, University of Southern California; MSTRA, Minnesota Study of Twins Reared Apart; MZ, monozygotic; DZ, dizygotic; *ns*, not significant (confidence interval spans 0); SFA, saturated fat; MUFA, monounsaturated fat; PUFA, polyunsaturated fat; CHO, carbohydrates; SC, simple carbohydrates; CC, complex carbohydrates.

*Heritability estimates are derived from the additive genetic effect value (a^2) calculated using structural equation modelling techniques as outlined by (Neale et al., 1992) unless otherwise specified. Twin pairs are same-sex. Ages are in years, unless otherwise specified. Energy intakes were calculated in kilocalories unless otherwise specified.

^aValues are age- and sex- adjusted. Energy intake calculated in kilojoules.

^bAge and sex-adjusted total caloric intake/meal heritability estimated only.

"Heritability estimates were calculated via Holzingers' H [(DZ variance-MZ variance)/DZ variance] or Falconer [2(rMZ-rDZ)] equations.

^dHeritability estimated using the LISREL method. Heritability per meal is provided in paratheses. ^cTwins reared apart study. Weight-adjusted heritability in parentheses (kilocalories/kg). Values are age- and gender- adjusted. ^fMZ within-pair exceeded between-pair mean square. ^gHeritability estimate derived from non-additive genetic effects (d²) due to correlation coefficient between MZ two times greater than DZ twins.

Table 2. S	election of th	vin studies	estimating ne	ritability of el	mpirically	derived food intake pat	terns (from Pallister et al., 2014).*	
			Dietary	T	Iwin			
Author	Sav	A a a	assessment	I WIN	pair	Diotory footor	Description	h ²
Author Faith at al	Sex Mala 8	Age	24 h rocall	MacArthur	222 MZ	Dietary factor	Eroquent intakes of legumes (peaputs and	<u>n</u> M: 70
(2008) ^a	Female $(n = 792)$,	(parents)	Longitudinal Study of Twins (US)	182 DZ	intake	peanut butter), and jam and jelly.	WI. 79
						Breakfast cereal and milk intake	Frequent intakes of milk and breakfast cereal.	ns
						Bread and butter intake	Frequent intakes of breads, butter and margarine.	<i>M</i> : 18 <i>F</i> : 20
						Adjusted fruit intake	Frequent intakes of non-citrus fruit, and fruit juice, punch and soda (reverse coded).	<i>M</i> : 26
						Adjusted red meat and pork intake	Frequent intakes of beef, pork, lamb and poultry (reverse coded).	<i>M</i> : 57
						Vegetable intake	Frequent intakes of deep-coloured vegetables and other vegetables.	ns
						Adjusted candy intake	Frequent intakes of candy, and sweets (reverse coded).	M: 41 F: 27
						Fish and lemon intake	Frequent intakes of fish and citrus fruit.	<i>M</i> : 12 <i>F</i> : 56
						High-salt snack food intake	Frequent intakes of high-salt snack foods.	<i>M</i> : 24
Keskitalo et	Male +	22-27	24-item FFQ	Finnish Twin	704 MZ	"Healthy foods"	Frequent intakes of fresh vegetables, fruits,	<i>M</i> : 49
<i>al.</i> (2007) ^b	female (<i>n</i> = 4018)	(x = 24.4)		Registry (Finland)	1490 DZ	·	cooked vegetables, berries, porridge, muesli, cereals, reduced-fat cheeses, rice or pasta, chicken yoghurt, and fish.	<i>F</i> : 54
						High-fat foods	Frequent intakes of fried foods, hamburgers, pizza, fried potatoes or French fries, creamy foods, and salty spaces	M: 44 F: 47
						Sweet foods	Frequent intakes of other sweets, chocolate, and sweet dessets	M: 42 E: 43
						Meat	High intakes of sausage and meat.	M: 39 F: 44
Teucher et	Female $(n = 3262)$	18-79 (x = 48 1)	131-item FFQ	Twins UK (UK)	498 MZ 1133 DZ	Fruit and vegetable	Frequent intakes of fruit, allium and cruciferous vegetables: low intakes of fried potatoes	43
un (2001)	((//)				High alcohol	Frequent intakes of beer, wine and allium vegetables; low intakes of high fibre breakfast cereals and fruit.	48
						Traditional English	Frequent intakes of fried fish and potatoes, meats, savoury pies and cruciferous vegetables.	41
						Dieting	Frequent intakes of low-fat dairy products, low- sugar soda; low intakes of butter and sweet baked products.	41
						Low meat	Frequent intakes of baked beans, pizza and soy foods; low intakes of meat, other fish and seafood and poultry	43
Breen <i>et al.</i> (2006) ^{b,c}	Male + female	4-5 (x = 4.4)	95-item modified FFQ	Twins Early Development	103 MZ 111 DZ	Vegetables	High liking of broccoli, cabbage, carrots, cauliflower, green beans, mushrooms, onions,	37
	(<i>n</i> = 428)		(parents)	Study (UK)		Desserts	parsnips, saiad greens and tomato. High liking of cream, cakes, pastries, fruit pie, sponge pudding, custard and dairy desserts.	20

						Meat and fish	High liking of beef, lamb, pork, chicken, bacon, fried fish, white fish and oily fish.	78
						Fruit	High liking of apples, bananas, citrus fruits, grapes, peaches, strawberries and fruit juice.	51
Gunderson <i>et al.</i> (2006) ^{b,d}	Female (<i>n</i> = 700)	30-90 (x̄ = 50)	100-item FFQ	Kaiser Permanente Twin Registry (US)	704 MZ 1490 DZ	'Healthy' dietary pattern	Frequent intakes of fish or chicken, carrots, tomatoes, salad, green or yellow vegetables, fruits, high fibre grains, rice and potatoes.	50
						'Unhealthy' dietary pattern	Frequent intakes of beef, pork, hot dogs, eggs, cheese, ice cream, butter, margarine, soda, and desserts.	ns
van den Bree <i>et al.</i> (1999) ^b	Male + female (<i>n</i> = 4640)	≥50	99-item semi- quantitative FFQ	Virginia Commonweal th University (US)	935 MZ 713 DZ	Less healthful	Frequent intakes of foods high in fat, salt and sugar.	33
				. ,		Healthful	Frequent intakes of a variety of vegetables, fruit, rice, yogurt, skim milk and dark bread.	33

FFQ, food frequency questionnaire; MZ, monozygotic; DZ, dizygotic; M, male; *ns*, not significant (confidence interval spans 0); F, female. *Heritability estimates are derived from the additive genetic effect value (a²) calculated using structural equation modelling techniques as outlined by (Neale et al., 1992) unless otherwise specified. Twin pairs are same-sex. Ages are in years, unless otherwise specified. ^aFood patterns identified through principal component analysis. ^bFood patterns identified through factor analysis. ^cHeritabilities derived from a modified 95-item food frequency questionnaire which asked food 'liking' as opposed to intake frequency.

^dUtilized a twins of mistaken zygosity approach.

					.			5			-	He	ritability	- /					
Author	Sex	Aqe	Method	Twin registry	Twin pair count	Veq	Potatoes	Fruit	Meat	Fish	Dairv	Eaas	Cereals	Legumes /nuts	Total	Sweets	Savourv	Fast food	Fats and oils
Pimpin <i>et</i> <i>al.</i> (2013) ^a	Male + female (<i>n</i> = 3605)	21 months	3-day diet diary	Gemini cohort (UK)	384 MZ 832 DZ	15	9	10	9 ns ^h		17	6 ns	9			5-15 ns ⁱ	4 ns		5 ns
Fildes <i>et al.</i> (2014) ^b	Male + Female (<i>n</i> = 2686)	3	114- item modified FFQ (parent)	Gemini cohort (UK)	458 MZ 872 DZ	54		53	48 ^h		27 ⁱ		32		29				
Hasselbalch <i>et al.</i> (2008) ^c	Male & female (<i>n</i> = 1212)	18-67 (x = 38)	247- item FFQ	Geminakar (Denmark)	600														
Males						24	68	-	34- 47 ^e	17	HF: 37 LF: 39	-	RG: 19 WG: 24			22 ns- 45 ^e			35- 48 ^e
Females						14 <i>n</i> s	28	-	29- 38 ⁱ	61	HF: 32 LF: 39	0 <i>ns</i>	RG: 12 <i>ns</i> WG: 20			23-27 ns ^e			9 <i>ns</i> - 42 ⁱ
Keskitalo <i>et</i> <i>al.</i> (2007)	Male + female (<i>n</i> = 4018)	22-27 (x̄ = 24.4)	24-item FFQ	Finnish Twin Registry (Finland)	704 MZ 1490 DZ														
Males	,					38-40 ^e	40-46 ^e	37- 51 ^e	22- 47 ⁱ	45	38- 48 ⁱ	30	42-49 ^e			23-55'	43	22- 55 ⁱ	
Females						48-50 ^e	38-44 ^e	44- 39 ^e	44- 49 ⁱ	44	37- 43 ⁱ	37	40-41 ^e			33-54 ⁱ	41	43- 54 ⁱ	
Teucher <i>et</i> <i>al.</i> (2007) ^d	Female (<i>n</i> = 3262)	18-79 (x̄ = 48.1)	131- item FFQ	Twins UK (UK)	498 MZ 1133 DZ	35-46 ^f	36 ⁹	40	29- 39 ^e	17 ns	HF: 24 LF: 36	29	RG: 8 WG: 29	30-32 ^e		30	7 ns	26	

Table 3. Selection of twin studies estimating heritability of food group intakes (from Pallister et al., 2014).*

FFQ, food frequency questionnaire; MZ, monozygotic; DZ, dizygotic; *ns*, not significant (confidence interval spans 0); HF, high fat; LF, low fat; RG, refined grain; WG, whole grain. *Heritability estimates are derived from the additive genetic effect value (a²) calculated using structural equation modelling techniques as outlined by (Neale et al., 1992) unless otherwise specified. Twin pairs are same-sex. '*ns*' indicates confidence interval spans 0. '-' indicates model including additive genetic effects not the best fit. Ages are in years, unless otherwise specified. Values presented were derived from single variables, unless otherwise specified.

^aHeritability analysis performed on age- and sex- adjusted residual scores.

^bHeritability analysis performed on age- and sex- adjusted residual scores.

^cHeritability analysis performed on food groups adjusted for total energy intake.

^dHeritability analysis performed on energy-adjusted residual scores.

^eTwo variables are included in the range.

^fFive variables are included in the range.

^gHertiability for root vegetable intake.

^hIncludes fish.

Three variables are included in the range.

¹Includes eggs.

							, .	,	Herita	bility			
Author	Sex	Age	Dietary assessment method	Twin registry	Twin pair count	Water	Alcohol	Soda	Diet soda	Milk	Coffee	Теа	Fruit juice
Pimpin <i>et</i> <i>al.</i> (2013) ^a	Male + female (<i>n</i> = 3605)	21 months	3-day diet diary	Gemini cohort (UK)	384 MZ 832 DZ	7		5 <i>ns</i> ^h		8			1 <i>n</i> s
de Castro <i>et</i> <i>al.</i> 1993 ^b	Male + female (<i>n</i> = 390)	x̄ = 38.8	7-day diet diary	Minnesota Twin Registry, (US)	109 MZ 86 DZ	80	73 ns	61	64	58 ns	73		-
Males Females				、 ,		76 ^f -	82 51	38 ⁱ 73	68 ⁱ 65 ⁱ	- 52	71 73		-
Hasselbalch et al. (2008) ^c	Male & female $(n = 1212)$	18-67 (x = 38)	247-item FFQ	Geminakar (Denmark)	600								
Males	(// = 1212)							26			63 ^j	63 ^j	36
Teucher <i>et</i> al. (2007) ^d	Female (<i>n</i> = 3262)	18-79 (x = 48.1)	131-item FFQ	Twins UK (UK)	498 MZ 1133 DZ		28	30			41	38	0 ns 0 ns
Hur <i>et al.</i> (1998) ^e	Male + female (<i>n</i> = 335)	18-77 (x = 42.4)	67-item FFQ	MSTRA (Multi- national)	66 MZ 51 DZ		38 ⁹	20 ns		33	29	46	25 ns

Table 4. Selection of twin studies estimating heritability of fluid intakes (from Pallister et al., 2014).*

FFQ, food frequency questionnaire; MSTRA, Minnesota Study of Twins Reared Apart; MZ, monozygotic; DZ, dizygotic; ns, not significant (confidence interval spans 0).

*Heritability estimates are derived from the additive genetic effect value (a²) calculated using structural equation modelling techniques as outlined by (Neale et al., 1992) unless otherwise specified. Twin pairs are same-sex. '-' indicates model including additive genetic effects not the best fit. Ages are in years, unless otherwise specified.

^aHeritability analysis performed on age- and sex- adjusted residual scores.

^bHeritability estimated using the LISREL method.

^cHeritability analysis performed on food groups adjusted for total energy intake.

^dHeritability analysis performed on energy-adjusted residual scores.

^eStudy used twins of mistaken zygosity approach. Heritability analysis performed on age- and gender- adjusted values.

^fSignificant dominant genetic effect

^gHeritability of alcohol type intake.

^hHeritability of other beverages group.

ⁱModel including additive genetic effects not best fitting model.

^jObtained from group 'other', includes coffee, tea and unspecified items.

Appendix B. Chapter 3 Appendices

Document 1. Food frequency questionnaire before 2007

CONFIDENTIAL

RHEUMATOLOGY DEPARTMENT OSTEOPOROSIS AND OSTEOARTHRITIS

TWIN RESEARCH



FOOD FREQUENCY QUESTIONNAIRE

This questionnaire asks for some background information about you especially what you eat.

Please answer every question. If you are uncertain about how to answer a question then do the best you can, but please do not leave a question blank.

Your answers will be treated as strictly confidential and will only be used for medical research.

PLEASE COMPLETE USING BLACK INK/PEN

STUDY NO.

DATE COMPLETED

YOUR DIET LAST YEAR

For each food there is an amount shown, either a 'medium serving' or a common household unit such as a slice or teaspoon. Please put a tick (\checkmark) in the box to indicate how often, on average, you have eaten the specified amount of each food during the past year.

EXAMPLES

For white bread the amount is one slice, so if you ate 4 or 5 slices a day, you should put a tick in the column headed '4-5' per day.

FOODS AND AMOUNTS	AVERAG	E USE	LAST	YEAR					
Bread & savoury biscuits (one slice or biscuit)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
White bread and rolls	e .							1	

For chips, the amount is a 'medium serving', so if you had a helping of chips twice a week you should put a tick in the column headed '2-4 per week'.

FOODS AND AMOUNTS	AVERAGE U	JSE LAST	YEAR					
Potatoes, Rice and Pasta (medium serving)	Never or less than once/month	1-3 Once per a ionth week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Chips			1					

For very seasonal fruits such as strawberries and raspberries you should estimate your average use when the fruits are in season, so if you ate strawberries or raspberries about once a week when they are in season you should put a tick in the column headed 'once a week'.

FOODS AND AMOUNTS	VERAGE	USE	LAST	YEAR					
Fruit (1 fruit medium serving)	ever or ss than nce/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6÷ per day
Strawberries, raspberries kiwi fruit			1						

Please estimate your average food use as best you can, and please answer every question - do not leave ANY lines blank. PLEASE PUT A TICK (/) ON EVERY LINE

FOODS AND AMOUNTS	AVERAGE L	JSE LAS	ST YEA	R					
MEAT AND FISH (medium serving)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Beef: roast, steak, mince, stew or casserole									
Beefburgers									
Pork: roast, chops, stew or slices									
Lamb: roast, chops or stew				-					
Chicken or other poultry eg. turkey									
Bacon									
Ham	-								
Corned beef, Spam, luncheon meats									
Sausages									
Savoury pies, eg. meat pie, pork pie, pasties, steak & kidney pie, sausage rolls									
Liver, liver paté, liver sausage									
Fried fish in batter, as in fish and chips									
Fish fingers, fish cakes									
Other white fish, fresh or frozen, eg. cod, haddock, plaice, sole, halibut									
Oily fish, fresh or canned, eg. mackerel, kippers, tuna, salmon, sardines, herring									
Shellfish, eg. crab, prawns, mussels									
Fish roe, taramasalata									
	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ pe da

Please check that you have a tick (\checkmark) on EVERY line

PLEASE PUT A TICK (1) ON EVERY LINE

FOODS AND AMOUNTS	AVERAGE U	JSE LAS	T YEA	R					
BREAD AND SAVOURY BISCUITS (one slice or biscuit)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
White bread and rolls									
Brown bread and rolls		13.00							
Wholemeal bread and rolls									
Cream crackers, cheese biscuits									
Crispbread, eg. Ryvita									
CEREALS (one bowl)									
Porridge, Readybrek									
Breakfast cereal such as cornflakes, muesli etc.	- 1								
POTATOES, RICE AND PASTA (medium s	erving)	1.1							
Boiled, mashed, instant or jacket potatoes									
Chips									
Roast potatoes									
Potato salad				100					
White rice									
Brown rice									
White or green pasta, eg. spaghetti, macaroni, noodles									
Wholemeal pasta									
Lasagne, moussaka									
Pizza									
	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day

Please check that you have a tick (\checkmark) on EVERY line

PLEASE PUT A TICK (1) ON EVERY LINE

belle and succession

FOODS AND AMOUNTS	AVERAGE L	JSE LAS	ST YEA	R					
DAIRY PRODUCTS AND FATS	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Single or sour cream (tablespoon)									
Double or clotted cream (tablespoon)									
Low fat yogurt, fromage frais (125g carton)									
Full fat or Greek yogurt (125g carton)									
Dairy desserts (125g carton)									
Cheese, eg. Cheddar, Brie, Edam (medium serving)									
Cottage cheese, low fat soft cheese (medium serving)									
Eggs as boiled, fried, scrambled, etc. (one)									
Quiche (medium serving)									
Low calorie, low fat salad cream(tablespoon)									
Salad cream, mayonnaise (tablespoon)									
French dressing (tablespoon)									
Other salad dressing (tablespoon)									
The following on bread or vegetables									
Butter (teaspoon)		2							
Block margarine, eg. Stork, Krona (teaspoon)									
Polyunsaturated margarine (tub), eg. Flora, sunflower (teaspoon)								-	
Other soft margarine,dairy spreads (tub), eg. Blue Band, Clover (teaspoon)									
Low fat spread (tub), eg. Outline, Gold (teaspoon)									
Very low fat spread (tub) (teaspoon)									
	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per dat

Please check that you have a tick (1/) on EVERY line

PLEASE PUT A TICK (/) ON EVERY LINE

FOODS AND AMOUNTS	AVERAGE	JSE LAS	ST YEA	R					
SWEETS AND SNACKS (medium serving)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Sweet biscuits, chocolate , eg. digestive (one)									
Sweet biscuits, plain, eg. Nice, ginger (one)									
Cakes eg. fruit, sponge, home baked									
Cakes eg. fruit, sponge, ready made									
Buns, pastries eg. scones, flapjacks, home baked									
Buns, pastries eg. croissants, doughnuts, ready made									
Fruit pies, tarts, crumbles, home baked									
Fruit pies, tarts, crumbles, ready made									
Sponge puddings, home baked									
Sponge puddings, ready made									
Milk puddings, eg. rice, custard, trifle							1		
Ice cream, choc ices									
Chocolates, single or squares			1						
Chocolate snack bars eg. Mars, Crunchie									
Sweets, toffees, mints									
Sugar added to tea, coffee, cereal (teaspoon)				1945					
Crisps or other packet snacks, eg. Wotsits									
Peanuts or other nuts									
SOUPS, SAUCES, AND SPREADS				10.45			Ser.		
Vegetable soups (bowl)									T
Meat soups (bowl)									
Sauces, eg. white sauce, cheese sauce, gravy (tablespoon)									
Tomato ketchup (tablespoon)									
Pickles, chutney (tablespoon)									
Marmite, Bovril (teaspoon)									-
Jam, marmalade, honey (teaspoon)									
Peanut butter (teaspoon)									
	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a dav	2-3 per dav	4-5 per dav	6+ pe da

Please check that you have a tick (✓) on EVERY line

PLEASE PUT A TICK (1) ON EVERY LINE

FOODS AND AMOUNTS	AVERAGE USE LAST YEAK										
DRINKS	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day		
Tea (cup)							2.6				
Coffee, instant or ground (cup)											
Coffee, decaffeinated (cup)											
Coffee whitener, eg. Coffee-mate (teaspoon)											
Cocoa, hot chocolate (cup)											
Horlicks, Ovaltine (cup)											
Wine (glass)											
Beer, lager or cider (half pint)											
Port, sherry, vermouth, liqueurs (glass)											
Spirits, eg. gin, brandy, whisky, vodka (single)											
Low calorie or diet fizzy soft drinks (glass)											
Fizzy soft drinks, eg. Coca cola, lemonade (glass)											
Pure fruit juice (100%) eg. orange, apple juice (glass)											
Fruit squash or cordial (glass)											
FRUIT (1 fruit or medium serving) For very seasonal fruits such as strawber	ries, please es	timate y	our ave	erage u	se wher	n the fru	lit is in	season			
Apples											
Pears											
Oranges, satsumas, mandarins					153						
Grapefruit				3.5							
Bananas											
Grapes				1.605							
Melon											
Peaches, plums, apricots											
Strawberries, raspberries, kiwi fruit											
Tinned fruit											
Dried fruit, eg. raisins, prunes											
	Never or less than	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per dav	4-5 per dav	6+ pe		

PLEASE PUT A TICK (1) ON EVERY LINE

FOODS AND AMOUNTS	AVERAGE USE LAST YEAR									
VEGETABLES Fresh, frozen or tinned (medium serving)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day	
Carrots										
Spinach										
Broccoli, spring greens, kale										
Brussels sprouts										
Cabbage										
Peas ,										
Green beans, broad beans, runner beans	2									
Marrow, courgettes	1									
Cauliflower										
Parsnips, turnips, swedes										
Leeks										
Onions										
Garlic				-						
Mushrooms										
Sweet peppers										
Beansprouts										
Green salad, lettuce, cucumber, celery										
Watercress										
Tomatoes										
Sweetcorn										
Beetroot										
Coleslaw				-				13.20		
Avocado		1997				-				
Baked beans										
Dried lentils, beans, peas										
Tofu , soya meat, TVP, Vegeburger										
	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day	

Please check that you have a tick (\checkmark) on EVERY line

Food	Lisual conving	Number of times
FOOU		Size eater each wee
*		
What type of milk did you	u most often use?	
Select one only Ful	I cream, silver	Semi-skimmed, red/white
	Skimmed/blue	Channel Islands, gold
	Dried milk	Soya
Other, specify		None _
How much milk did you o	drink each day, including mi	ilk with tea, coffee, cereals etc?
	None	Three quarters of a pint
Qu	uarter of a pint	One pint
	Half a pint	More than one pint
Did you usually eat breal	kfast cereal (excluding porrid	ge and Ready Brek mentioned earlier)? Yes No _
Did you usually eat breat If yes, which brand and t	kfast cereal (excluding porrid type of breakfast cereal, inc	ge and Ready Brek mentioned earlier)? Yes No Ves No Cluding muesli, did you usually eat?
Did you usually eat breat If yes, which brand and t List the one or two type Brand	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ	ge and Ready Brek mentioned earlier)? Yes No _ cluding muesli, did you usually eat?
Did you usually eat breat If yes, which brand and t List the one or two type Brand	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ	ge and Ready Brek mentioned earlier)? Yes No cluding muesli, did you usually eat?
Did you usually eat breat If yes, which brand and t List the one or two type Brand	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ	ge and Ready Brek mentioned earlier)? Yes No _ cluding muesli, did you usually eat?
Did you usually eat breat If yes, which brand and i List the one or two type Brand	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ	ge and Ready Brek mentioned earlier)? Yes No Cluding muesli, did you usually eat?
Did you usually eat break If yes, which brand and t List the one or two type Brand What kind of fat did you	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ	ge and Ready Brek mentioned earlier)? Yes No cluding muesli, did you usually eat? De Desting, grilling etc?
Did you usually eat break If yes, which brand and t List the one or two type Brand What kind of fat did you Select one only	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ most often use for frying, ro Butter	ge and Ready Brek mentioned earlier)? Yes No cluding muesli, did you usually eat? De De Dasting, grilling etc? Solid vegetable fat
Did you usually eat break If yes, which brand and the List the one or two type Brand What kind of fat did you Select one only	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ most often use for frying, ro Butter Lard/dripping	ge and Ready Brek mentioned earlier)? Yes No cluding muesli, did you usually eat? De Dasting, grilling etc? Solid vegetable fat Margarine
Did you usually eat break If yes, which brand and t List the one or two type Brand What kind of fat did you Select one only	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ most often use for frying, ro Butter Lard/dripping Vegetable oil	ge and Ready Brek mentioned earlier)? Yes No cluding muesli, did you usually eat? De Desting, grilling etc? Solid vegetable fat Margarine None
Did you usually eat break If yes, which brand and the List the one or two type Brand What kind of fat did you Select one only If you used vegetable of	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ most often use for frying, ro Butter Lard/dripping Vegetable oil bil, please give type eg. cor	ge and Ready Brek mentioned earlier)? Yes No cluding muesli, did you usually eat? De Des Des Des Des Des Des Des Des Des
Did you usually eat break If yes, which brand and the List the one or two type Brand What kind of fat did you Select one only If you used vegetable of What kind of fat did you	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ most often use for frying, ro Butter Lard/dripping Vegetable oil Dil, please give type eg. cor most often use for baking c	ge and Ready Brek mentioned earlier)? Yes No cluding muesli, did you usually eat? De Dasting, grilling etc? Solid vegetable fat Margarine None rn, sunflower
Did you usually eat break If yes, which brand and the List the one or two type Brand What kind of fat did you Select one only If you used vegetable of What kind of fat did you Select one only	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ most often use for frying, ro Butter Lard/dripping Vegetable oil Dil, please give type eg. cor most often use for baking c Butter	ge and Ready Brek mentioned earlier)? Yes No cluding muesli, did you usually eat? De Desting, grilling etc? Solid vegetable fat Margarine None rn, sunflower cakes etc? Solid vegetable fat
Did you usually eat break If yes, which brand and the List the one or two type Brand What kind of fat did you Select one only If you used vegetable of What kind of fat did you Select one only	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ most often use for frying, ro Butter Lard/dripping Vegetable oil bil, please give type eg. cor most often use for baking c Butter Lard/dripping	ge and Ready Brek mentioned earlier)? Yes No Cluding muesli, did you usually eat? De De De De De De De De De De De De De
Did you usually eat break If yes, which brand and the List the one or two type Brand What kind of fat did you Select one only If you used vegetable of What kind of fat did you Select one only	kfast cereal (excluding porrid type of breakfast cereal, inc es most often used Typ 	ge and Ready Brek mentioned earlier)? Yes No Cluding muesli, did you usually eat? De De De De De De De De De De De De De

8.	How often did you eat food that was fried at home?	
	Daily 1-3 times a week	4-6 times a week
	Less than once a week	Never
9.	How often did you eat fried food away from home?	
	Daily 1-3 times a week	4-6 times a week
	Less than once a week	Never
10.	What did you do with the visible fat on your meat?	
	Ate most of the fat	Ate as little as possible
	Ate some of the fat	Did not eat meat
11.	How often did you eat grilled or roast meat?	times a week
12.	How well cooked did you usually have grilled or roast meat?	
	Well done /dark brown	Lightly cooked/rare
	Medium	Did not eat meat
13.	How often did you add salt to food while cooking?	
	Always	Rarely
	Usually	Never
	Sometimes	
14.	How often did you add salt to any food at the table?	
	Always	Rarely
	Usually	Never
	Sometimes	
15.	Did you regularly use a salt substitute (eg LoSalt)?	Yes No
	If yes, which brand?	
	10	

Have you taken any vitamins,	minerals, fish	h oils, fibre or of	ther food supplements	during the
past year?				

YES NO DON'T KNOW

If yes, please complete the table below. If you have taken more than 5 types of supplement please put the most frequently consumed brands first.

Vitamin supplements	Tic	Average frequency Tick one box per line to show how often on average you consumed supplements									
Name and brand Please list full name, brand and strength	Dose Please state number of pills, capsules or teaspoons consumed	Never or less than once a month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per oay	
		-	+								

Thank you for your help

Document 2: Food frequency questionnaire after 2007

Page 1

Study Number:



Twin Research & Genetic Epidemiology Unit

King's College London, St Thomas' Campus, Lambeth Palace Road, London SE1 7EH Tel: 0207 188 5555, Fax: 0207 188 6761 Email: twinrecruitment@gstt.nhs.uk Website: www.twinsuk.ac.uk

June 2007

Dear Twin,

We are excited to send you a new questionnaire about food and nutrition developed with our colleagues at the University of East Anglia in Norwich.

The aim of this important study is to identify the genetic variations that influence our individual response to the food we eat, including vitamins and minerals. We are also including questions about your general health, which may be affected by your diet.

You may have answered some of the questions several years ago, but we need to keep our information as up to date as possible, so please do complete all the questions asked. We would like to build up a full profile of how diet may be related to health status. Please complete the questionnaire using a black pen. If you have any queries regarding this questionnaire please contact Dr Paula Skidmore on 01603 591270.

Thank you very much for your time and your continued support. Please return your completed questionnaire in the pre-paid envelope enclosed.

Yours sincerely,

Professor Tim Spector Director, Twin Research & Genetic Epidemiology Unit





YOUR DIET LAST YEAR

Page 2

For each food there is an amount shown, either a "medium serving" or an example of a serving such as "a slice" or "a teaspoon". Please put a cross (*) in the box to indicate how often, **on average**, you have eaten a specified amount of food over the last 12 months. If you complete a box <u>incorrectly</u> please fill the wrong one. e.g. and put a cross in the correct box.

The rest of this page shows examples of how to fill in the questionnaire.

These questions are all about the food you have eaten over the last year.



For white bread the serving size is one slice or roll, so if you ate 2 to 4 slices a week on average, over the last 12 months, you should put a cross in the column headed "2-4 per week".

neaded 2-4 per week .										
FOODS AND AMOUNTS AVERAGE USE IN THE LAST TWELVE MONTHS										
BREAD (one slice or roll)		Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
White bread and rol	lls				×	2		9		xample o
EXAMPLE 2 For chips, the serving size is a small side plate so if you ate a serving of chips twice a week, over the last 12 months, you should put a cross in the column headed "2-4 per week".										
FOODS AND AMO	UNTS	AVER	AGE U	SE IN T	THE LA	ST TW	ELVE	MONT	HS	
POTATOES (one small side plate Chips	e)	Never or less than once/month	1-3 per month	Once a week	× 2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
EXAMPLE 3	For very seasor your average in a week in summ UNTS	nal fruits take in s ner you s AVER	such as ummer should p AGE U	s strawl , so if y out a cr SE IN 1	berries ou ate oss in t THE LA	and ra strawb he colu	spberri erries o imn he /ELVE	es you or raspt aded "o MONT	should perries a once a v	estimate about on week".
FRUIT (1 fruit or handful)		Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Strawberries, raspb	erries, kiwi fruit			×						
l										

	Page 3			Stud	ly Numb	er:					
STA	START OF QUESTIONNAIRE										
1a.	On average do you think you eat the s less than someone of your own age as Please put a cross (*) in only one b	ame, m nd sex? ox.	ore or		Sar	ne	1	More		Less	
1b.	On an average day how many portions	s of fruit	do you	eat?							
1c.	On an average day how many portions	s of veg	etables	do you	u eat?						
Please estimate your average food use as best you can, and please answer every question - do not leave ANY lines blank. PLEASE PUT A CROSS (*) ON EVERY LINE											
FOC	DS AND AMOUNTS	AVERA	GE US	E IN TI	HELAS	T TWI	ELVE	NONTH	IS		
DAII	RY PRODUCTS AND FATS	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day	
Sing	le or sour cream (tablespoon)										
Dout	ble or clotted cream (tablespoon)										
Low	fat yoghurt, fromage frais (small pot)										
Full f	at or Greek yoghurt (small pot)										
Dairy mou	desserts (small pot) e.g. chocolate sse, cream caramels										
Chee (mat	ese, e.g. cheddar, brie, edam chbox size)										
Low (mat	fat cheese e.g. reduced fat cheddar chbox size)										
Cotta (2 ta	age cheese, low fat soft cheese blespoons)										
Eggs	as boiled, fried, scrambled, etc. (one)										
Quic	he (slice)										
Low	calorie, low fat salad cream (tablespoon)										
Full f	at salad cream, mayonnaise (tablespoon)										
Fren	ch dressing (tablespoon)										
Othe	r salad dressing (tablespoon)										

Please check that you have a cross (×) on EVERY line

_	
	3280 R04501 118
	Page 4
	I ago T

Study	Number:
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day

6+ per

FOODS AND AMOUNTS AVERAGE USE IN THE LAST TWELVE MONTHS Never or less than once/month The following list refers to **DAIRY PRODUCTS AND FATS** month Once a week week week Once a day 2-3 per day 4-5 per day that you put on bread or cooked 5-6 per v 2-4 per v vegetables 1-3 per Butter (teaspoon) Reduced fat butter (teaspoon) Block margarine, e.g. Stork, Krona (teaspoon) Polyunsaturated margarine, e.g. Flora, sunflower (teaspoon) Olive oil spread (teaspoon) Other soft margarine, dairy spreads, e.g. Blue Band, Clover (teaspoon) Low fat spread, e.g. Outline, Gold (teaspoon) Very low fat spread (teaspoon) e.g. Diet Flora Cholesterol lowering fat spreads e.g. Benecol (teaspoon)

FRUIT - For seasonal fruits marked*, please estimate your average use in summer

Apples (1 fruit)					
Pears (1 fruit)					
Oranges, satsumas, mandarins (1 fruit)					
Grapefruit (half)					
Bananas (1fruit)					
Grapes (handful)					
Melon (1 slice)					
* Peaches, plums, apricots (1 fruit)					
* Strawberries, raspberries, other berries, kiwi fruit (one fruit or handful)					
Tinned fruit (handful)					
Dried fruit, e.g. raisins, prunes (heaped tablespoon)					

Please check that you have a cross (*) on EVERY line
Page 5			St	udy Nur	nber:				
FOODS AND AMOUNTS	AVEF	RAGE L	JSE IN	THE L	AST T	WELVE	MON	THS	
VEGETABLES Fresh, frozen or tinned (handful)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Carrots Spinach Broccoli, spring green, kale									
Brussel sprouts Cabbage									
reas Green beans, broad beans, runner beans Baked beans									
Marrow, courgettes Cauliflower Parsnips, turnips, swedes									
Leeks									
Garlic (clove) Mushrooms Sweet peppers									
Beansprouts Green salad, lettuce, cucumber, celery									
Watercress Tomatoes									
Beetroot Coleslaw									
Avocado Pulses e.g. lentils, beans, peas Meat substitutes e.g. tofu, soyameat.									



Study Number:

FOODS AND AMOUNTS

Page 6

AVERAGE USE IN THE LAST TWELVE MONTHS

MEAT AND FISH (half a small side plate)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Beef: roast, steak, mince, stew or casserole									
Beefburgers									
Pork: roast, chops or stew									
Lamb: roast, chops or stew									
Chicken or other poultry e.g. turkey									
Bacon or gammon									
Ham, cured meats & chorizo									
Corned Beef, Spam, luncheon meats									
Sausages									
Savoury pies, e.g. meat pie, pork pie, pasties, steak & kidney pie, sausage rolls									
Liver, liver pate, liver sausage									
Fried fish in batter, as in fish and chips									
Fish fingers, fish cakes & breaded fish									
Other white fish, fresh or frozen, e.g. cod, plaice, sole, haddock, halibut									
Oily fish, fresh or canned, e.g. tuna, mackerel, kippers, salmon, sardines, herring									
Shellfish, e.g. crab, prawns, mussels									
Fish roe, taramasalata									

Please check that you have a cross (×) on EVERY line



Study Number:

FOODS AND AMOUNTS

AVERAGE USE IN THE LAST TWELVE MONTHS

BREAD AND SAVOURY BISCUITS (one slice, or roll, or biscuit)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
White bread/rolls		\square			\square				
Brown bread/rolls					\Box				
Wholemeal & granary bread/rolls		Π	\square	\square	\square	\square	Π		
Cream crackers, savoury biscuits									
Crispbread, e.g. Ryvita									
Naan, poppadoms, flour tortillas									

SOUPS, SAUCES AND SPREADS

Vegetable soups (bowl)	
Meat soups (bowl) (to include meat and vegetable soups)	
Sauces, e.g. white sauce, cheese sauce, gravy (tablespoon)	
Tomato ketchup (tablespoon)	
Pickles, chutney (tablespoon)	
Marmite, Bovril (teaspoon)	
Jam, marmalade, honey (teaspoon)	
Peanut butter (teaspoon)	

Please check that you have a cross (*) on EVERY line

Page 8

Study Number:



8

FOODS AND AMOUNTS

AVERAGE USE IN THE LAST TWELVE MONTHS

SWEETS AND SNACKS (medium serving)	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Cakes e.g. fruit, sponge, <u>home baked</u>									
Cakes e.g. fruit, sponge, ready made									
Buns, pastries e.g. scones, flapjacks, croissants, doughnuts, <u>home baked</u>									
Buns, pastries e.g. scones, flapjacks, croissants, doughnuts, <u>ready made</u>									
Fruit pies, tarts, crumbles, <u>home baked</u>									
Fruit pies, tarts, crumbles, ready made									
Sponge puddings, home baked									
Sponge puddings, ready made									
Milk puddings e.g. rice, custard, trifle									
Ice cream, choc ices									
Sweet biscuits, chocolate, e.g. digestive (one)									
Sweet biscuits, plain, e.g. Nice, ginger (one)									
Reduced fat biscuits e.g. Go Ahead, Highlights (one small packet or one small bar/biscuit)									
Cereal bars (one)									
White or milk chocolates, single or squares (one)									
Dark chocolates, single or squares (one)									
Chocolate snack bars e.g. Mars, Crunchie (one)									
Sweets, toffees, mints (small packet)									
Sugar added to tea, coffee, cereal (teaspoon)									

Please check that you have a cross (*) on EVERY line



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FOODS AND AMOUNTS

Study Number:



AVERAGE USE IN THE LAST TWELVE MONTHS

SWEETS AND SNACKS	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Crisps or other packet snacks, e.g. Wotsits (one packet)									
Salted nuts e.g. peanuts, cashews (handful)									
Unsalted nuts, e.g. brazil, walnuts (handful)									
Seeds e.g. Sunflower, pumpkin (tablespoon)									
CEREAL (one bowl)									
Porridge, Readybreak, oats									
Breakfast cereal e.g. Cornflakes, Rice Krispies									
Sugar topped cereals e.g. Frosties									
Muesli									
High Fibre cereals e.g. Branflakes, All Bran, Fruit and Fibre									
POTATOES, RICE AND PASTA (one small side plate)									
Boiled, mashed, instant or one jacket potato									
Chips, roast potatoes									
Potato salad									
White rice									
Brown rice									
White or green pasta, e.g. spaghetti, macaroni, noodles									
Wholemeal pasta									
Lasagne, moussaka									
Pizza (one slice)									
Please check that you h	iave a) Cros	s (*)	on E	EVER	IY lin)@		



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Study Number:



8

FOODS AND AMOUNTS

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AVERAGE USE IN THE LAST TWELVE MONTHS

DRINKS	Never or less than once/month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per day	4-5 per day	6+ per day
Tea (cup)									
Green tea (cup)									
Fruit tea (cup)									
Coffee, instant or ground (cup)									
Coffee, decaffeinated (cup)									
Coffee whitener, e.g. Coffee-mate (teaspoon)									
Cocoa, hot chocolate (cup)									
Low fat hot chocolate (cup)									
Horlicks, Ovaltine (cup)									
White wine (small glass)									
Red wine (small glass)									
Beer, lager or cider (half pint)									
Port, sherry, vermouth, liqueurs (pub measure)									
Spirits, e.g. gin, brandy, whisky, vodka (pub measure)									
Low calorie or diet fizzy soft drinks (cup)									
Fizzy soft drinks, e.g. Coca Cola, lemonade (cup)									
Pure fruit juice (100%) e.g. orange, apple juice (cup)									
Fruit squash or cordial (cup)									
Smoothies (cup)									

Please check that you have a cross (*) on EVERY line



Page 11	Study Number	:						
OUR DIET LAST YEAR, continu	ed	· · · · · · · · · · · · · · · · · · ·						
Are there any OTHER foods whi	ch you ate more than once a week o	over the last year?						
No 🔄 🖨 GO TO question 3	Yes If yes, please list b	oelow						
Food	Usual serving size	Number of times eaten each week						
Have you drunk milk or a milk su	bstitute over the past year? (includir Yes	ng in hot drinks) nn 4						
What type of milk or milk substitu	ute did you <u>most often use</u> over the l	last year?						
Select one only Full c Skimr Dried Other (please	ream	Semi-skimmed						
About how much milk did you dr	ink each day, including milk with tea	, coffee, cereals etc?						
Less than a quarter of a pint Half a pint Three quarters of a pint	One Pint More than or	ne pint						
Did you usually eat breakfast ce mentioned earlier)?	real in the last year (excluding Porric	dge and Ready Brek						
No 🔜 🖙 GO TO question 7	Yes If yes, which brancereal, including r	nd and type of breakfast muesli, did you usually eat?						
List the one or two types most Brand e.g. Kellogg's	t often used Type e.g. cornfl	akes						
Do you use oil or fat for frying, ro	pasting, grilling etc?							
No ⇔ GO TO question 9) Yes SO TO ques	tion 8						
i								

E P	Page 12		Study Number:									
8.	What kind of fat did you Butter	most often use for fryin Olive oil Solid vegetable fat	ng, roasting, grilling etc? Sele Margarine Low fat spray	ct one only Vegetable oil								
	If you used vegetable of please give type e.g. co	bil or other oil rn, sunflower, sesame										
9.	What kind of fat did you I don't bake	most often use for baki Lard/dripping	ing cakes etc? Select one or Solid vegetable fat	None								
	If you used margarine, please give name or type e.g. Flora, Stork											
10.	How often did you eat for Never	ood that was fried at ho Less than Daily	me? once a week	1-3 times a week								
11.	How often did you eat fr Never	ied food away from hor Less than Daily	ne? once a week	1-3 times a week								
12.	Have you eaten meat in No ⇔ GO TO que	the last year? stion 17 Yes	GO TO question 13									
13.	What did you do with the Did not eat meat with vis Ate as little as possible	e visible fat on your me sible fat	at? Ate some of the Ate most of the f	fat								
14.	Did you eat roast/grilled	meat over the past yea	ar?									
	No⇒ GO TO ques	stion 17 Yes	⇒ GO TO question 15									
15.	How often did you eat g	rilled or roast meat?										
	Daily		1-3 times a week]								

	Page 13	Stu	dy Number:	
16.	How well cooked did you usually have g	rilled or roast meat?	,	
	Well done / dark brown		Lightly coo	ked / rare
17.	How often did you add salt to food while	cooking?		
	Never	Sometimes		Always
18.	How often did you add salt to any food a	at the table?		
	Never Rarely	Sometimes		Always
19.	Do you regularly use a salt substitute (e	.g. LoSalt)?		
	No Yes	If yes, which	h brand?	
20.	We would like to summarise what you h average, how many times a week did yo	ave told us. During t ou eat the following f	he course of la oods?	ast year, on
	Food Type	Times	/week	Portion size
	Vegetables (excluding potatoes but including baked beans)			medium serving
	Salads			medium serving
	Fruit and fruit products (not including fru	it juice)		medium serving or 1 fruit
	Oily fish and fish products			medium serving
	Non oily fish and fish products			medium serving
	Meat, meat products and meat dishes (including bacon, ham and chicken)			medium serving
21.	Have you taken any vitamins, minerals, the past year?	fish oils, fibre or oth	er food supple	ments during
	No ⇔ GO TO section on HEAL on page 15	.TH Yes 🗌 ⇔	GO TO ques	tion 22







22. Please list any vitamins, minerals, fish oils, fibre or other food supplements taken during the past year. If you have taken more than 6 types of supplement please list the 6 most frequently consumed brands first.

Vitamin supplements			Cros on a	Avera s one box	age fre k per line ou consur	to show h	y ow often lements			
Name and brand Please list full name and brand	Number Please state number of pills, capsules or teaspoons consumed	Never or less than once a month	1-3 per month	Once a week	2-4 per week	5-6 per week	Once a day	2-3 per say	4-5 per day	6+ per day
	1 - 9									
	1 - 9									
	1 - 9									
	1 - 9									
	1 - 9									

Appendix C. Chapter 4 Appendices

Table 1. Study	population	characteristics
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	Whole populat	ion (<i>n</i> =3559)	Discordant monozygotic twins		
Variable	Mean (SD)	Range	<i>n</i> pairs	Mean	Range
Age (years)	55.3 (13.4)	18 - 84			_
Body mass index (kg/m ²)	26.1 (4.9)	13.6 - 50.1			
Intakes (servings/week)	. ,				
Allium vegetables	4.5 (4.2)	0 - 84.5	174	7.1 (5.6)	0 - 37
Apples/pears	4.7 (4.9)	0 - 74	206	7.0 (6.2)	0 - 36
Avocado	0.4 (0.9)	0 - 18	129	1.6 (2)	0 - 18
Baked beans	0.9 (1.0)	0 - 7	149	1.8 (1.6)	0 - 7
Bananas	39(37)	0 - 42	210	5 3 (5)	0 - 32
Beef burgers	0.1 (0.3)	0-3	172	0.3(0.4)	0 - 3
Beer	0.7(2.4)	0 - 42	63	35(56)	0 - 32
Berries	0.7(2.4)	0 - 14	137	21(27)	0 - 14
Black tea	18 0 (13 8)	0 - 42	240	18/	0 - 42
Butter	35(61)	0 - 50	173	85(80)	0 = 42
Cheese	3.3(0.1)	0-36	109	5.0 (4.8)	0 - 42
Cheese	3.1(3.1)	0 - 30	190	5.0 (4.0) 9.0 (40.7)	0 - 30
Chocolate	3.8 (6.1)	0 - 84	153	8.9 (10.7)	0 - 84
Citrus Iruit	3.0 (4.3)	0 - 45	167	0.2 (0.0)	0 - 36
Contee	12.3 (12.4)	0 - 84	252	17.4	0 - 84
Cooked potatoes	3.4 (2.6)	0 - 42	207	4.1 (3.3)	0 - 32
Cream	0.5 (1.3)	0 - 36	70	2.1 (2.1)	0 - 8
Crispbread	1.2 (3.4)	0 - 42	85	6.1 (8.6)	0 - 42
Cruciferous vegetables	6.1 (4.7)	0 - 98	219	8.6 (5.7)	0 - 35
Dairy desserts	0.2 (0.7)	0 - 18	94	0.9 (1.8)	0 - 18
Eggs	1.4 (1.6)	0 - 18	209	2.6 (2.5)	0 - 18
Fried fish	0.5 (0.7)	0 - 11	195	0.9 (1)	0 - 7.5
Fried potatoes	1.4 (1.4)	0 - 32	184	2.2 (1.7)	0 - 12
Fruit juice	3.4 (4.3)	0 - 42	221	5.8 (6)	0 - 42
Grapes	2.0 (2.9)	0 - 42	161	4.3 (4.7)	0 - 32
Green leafy vegetables	4.7 (4.4)	0 - 53	186	8.0 (7)	0 - 43
Herbal tea	1.6 (5.2)	0 - 64	117	9.5 (10.3)	0 - 50
High fat milk	1.0 (4.0)	0 - 45.5	59	7.4 (10.4)	0 - 45.5
High fibre breakfast	2.7 (3.5)	0 - 42	220	4.1 (4.1)	0 - 32.5
High sugar drinks	2.5 (5.6)	0 - 60	109	9.1 (10.2)	0 - 49
Ice cream	0.7 (1.2)	0 - 18	132	2.0 (1.8)	0 - 7
Lasagne	0.3 (0.4)	0 - 3	307	0.4 (0.4)	0 - 3
Legumes	4.3 (3.3)	0 - 43	230	5.8 (4.2)	0 - 32
Low fat milk	3.3 (2.4)	0 - 10.5	228	3.6 (3)	0 - 10.5
Low fibre breakfast cereals	1.6 (2.9)	0 - 42	224	3.4 (3.9)	0 - 42
Low fat spread	1.4 (4.5)	0 - 74	90	6.8 (8.5)	0 - 42
Malt drinks	0.3 (1.4)	0 - 32	66	2.9 (3.1)	0 - 18
Margarine	2.3 (4.9)	0 - 64	186	6.5 (7.8)	0 - 42.5
Marrow	0.7 (1.3)	0 - 32	130	2.1 (1.8)	0 - 7
Meat	2.4 (2.0)	0 - 16.5	222	3.3 (2.4)	0 - 11
Melon	0.9 (1.7)	0 - 32	150	2.8 (3.4)	0 - 32
Mushrooms	1.5 (1.7)	0 - 32	258	2.4 (1.8)	0 - 7
Nuts	17(32)	0 - 43 5	144	5.0 (5.3)	0 - 41 5
Oily fish	1 1 (1 2)	0 - 18	217	21(17)	0 - 18
Other fish/seafood	12(12)	0 - 16 5	171	21(17)	0-95
Peaches	0.7(1.2)	0 - 14	141	1.8 (2.2)	0 - 14
Pizza	0.5(0.7)	0-8	199	0.8 (0.9)	0-55
Polyunsaturated margarine	3.1 (6.0)	0 - 42	143	9.3 (10)	0 - 42
Porridae	1 4 (2 4)	$0 = \pm 2$	215	3.0(27)	$0 = \frac{1}{2}$
Pointage	1.4(2.4)	0-32	210	3.0(2.7)	0-7
Processed maste	1.3 (1.3)	0 60	100	2.2 (1.4) 1 2 (1 2)	0 22
Processed meats	1.0(3.1)	0 - 00	190	4.3 (4.2)	0 - 32
Frocessed fruit	2.8(2.3)	0 - 18	220	3.9 (2.8)	0 - 15.5
	4.0 (3.7)	0 - 96	202	0.3 (4.5)	0 - 28
righ fat salad dressing	1.6 (2.2)	0 - 25	201	3.1 (3.3)	0 - 25
Low fat salad dressing	0.7 (1.4)	0 - 18	140	2.3 (2.4)	0 - 18

	Whole populat	tion (n-2550)	Discor	dant monozva	otic twine
		uon (<i>n</i> =3559)	DISCOL		
Variable	Mean (SD)	Range	<i>n</i> pairs	Mean	Range
Savoury pies	0.3 (0.5)	0 - 7	163	0.5 (0.8)	0 - 7
Savoury snacks	2.1 (3.0)	0 - 35	170	4.5 (5.3)	0 - 35
Seasonings	3.6 (3.7)	0 - 84.5	195	5.6 (6.6)	0 - 84.5
Low sugar soda	1.8 (4.8)	0 - 42	92	7.9 (9.2)	0 - 42
Soup	1.1 (1.9)	0 - 64	149	2.9 (2.6)	0 - 14
Soy/other milk	0.1 (0.8)	0 - 10.5	68	1.7 (1.9)	0 - 10.5
Soy foods	0.2 (1.0)	0 - 18	60	2.2 (3.2)	0 - 18
Spirits/liquor	1.2 (3.2)	0 - 50	97	5.1 (7)	0 - 42.5
Sweet baked products	7.9 (8.9)	0 - 85	155	15.5	0 - 76.5
Sweetcorn	0.9 (1.3)	0 - 18	187	2.1 (1.9)	0 - 18
Sweet peppers	1.4 (1.8)	0 - 32	245	2.5 (2.1)	0 - 18
Confectionary/jam	8.1 (11.9)	0 - 84.5	149	18.2	0 - 84
Tomatoes	4.0 (3.6)	0 - 42	175	6.5 (6.2)	0 - 42
White/brown bread, refined	8.3 (8.5)	0 - 70	206	12.8	0 - 43.5
Wholemeal bread/grains	4.9 (6.7)	0 - 44	179	10.3 (9.2)	0 - 43
Wine	3.9 (6.1)	0 - 64	111	12.2	0 - 43
Yoghurt	2.9 (3.7)	0 - 47.5	180	5.3 (5.6)	0 - 47.5

Table 1. Study population characteristics

Food group	FEQ Variable Name	FFQ Description	Included	Included
Allium vegetables	Garlic	Garlic (clove)	TRUE	TRUE
Allium vegetables	Leeks	Leeks	TRUE	TRUE
Allium vegetables	Onions	Onions	TRUE	TRUE
Apples/pears	Apples	Apples (1 fruit)	TRUE	TRUE
Apples/pears	Pears	Pears (1 fruit)	TRUE	TRUE
Avocado	Avocado	Avocado	TRUE	TRUE
Baked beans	BakedBeans	Baked beans	TRUE	TRUE
Bananas	Bananas	Bananas (1fruit)	TRUE	TRUE
Beef burgers	MFburgers	Beefburgers	TRUE	TRUE
Beer	Beer	Beer, lager or cider (half pint)	TRUE	TRUE
Berries	Strawberries	Strawberries, raspberries, other berries, kiwi fruit (one fruit or handful)	TRUE	TRUE
Black tea	Теа	Tea (cup)	TRUE	TRUE
Butter	RedFatButter	Reduced fat butter (teaspoon)	FALSE	TRUE
Butter	Butter	Butter (teaspoon)	TRUE	TRUE
Cheese	DairyLFcheese	Low fat cheese e.g. reduced fat cheddar (matchbox size)	FALSE	TRUE
Cheese	Dairycheese	Cheese, e.g. cheddar, brie, edam (matchbox size)	TRUE	TRUE
Cheese	DairyCottageCheese	Cottage cheese, low fat soft cheese (2 tablespoons)	TRUE	TRUE
Chocolate	ChocsDark	Dark chocolates, single or squares (one)	FALSE	TRUE
Chocolate	ChocsMilk	White or milk chocolates, single or squares (one)	FALSE	TRUE
Chocolate	LowFatHotChoc	Low fat hot chocolate (cup)	FALSE	IRUE
Chocolate	Cocoa	Cocoa, hot chocolate (cup)	TRUE	TRUE
Chocolate	Chocolatebar	Chocolate snack bars e.g. Mars, Crunchie (one)	TRUE	TRUE
Chocolate	Chocolates	Chocolate, fancy and filled	TRUE	FALSE
Citrus fruit	Grapefruit	Grapefruit (half)	TRUE	TRUE
Citrus fruit	Oranges	Oranges, satsumas, mandarins (1 fruit)	TRUE	TRUE
Coffee	Coffee	Coffee, instant or ground (cup)	TRUE	TRUE
Coffee	CoffeeDecaffeinated	Coffee, decaffeinated (cup)	TRUE	TRUE
Confectionary/ja m	Jam	Jam, marmalade, honey (teaspoon)	TRUE	TRUE
Confectionary/ja m	SugarAdded	Sugar added to tea, coffee, cereal (teaspoon)	TRUE	TRUE
Confectionary/ja m	SweetsToffees	Sweets, toffees, mints (small packet)	TRUE	TRUE
Cooked potatoes	Boiledpotato	Boiled, mashed, instant or one jacket potato	TRUE	TRUE
Cream	Dairydouble	Double or clotted cream (tablespoon)	TRUE	TRUE
Cream	Dairysingle	Single or sour cream (tablespoon)	TRUE	TRUE
Crispbread	Crispbread	Crispbread, e.g. Ryvita	TRUE	TRUE
Cruciferous vegetables	Broccoli	Broccoli, spring green, kale	TRUE	TRUE
Cruciferous vegetables	BrusselsSprouts	Brussel sprouts	TRUE	TRUE

			Included	Included
Food group	FFQ Variable Name	FFQ Description	FFQ1	FFQ2
Cruciferous vegetables	Cabbage	Cabbage	TRUE	TRUE
Cruciferous vegetables	Cauliflower	Cauliflower	TRUE	TRUE
Cruciferous vegetables	Coleslaw	Coleslaw	TRUE	TRUE
Dairy desserts	DairyDesserts	Dairy desserts (small pot) e.g. chocolate mousse, cream caramels	TRUE	TRUE
Eggs	Eggs	Eggs as boiled, fried, scrambled, etc. (one)	TRUE	TRUE
Fried fish	MFFishFingers	Fish fingers, fish cakes & breaded fish	TRUE	TRUE
Fried fish	MFFriedfish	Fried fish in batter, as in fish and chips	TRUE	TRUE
Fried potatoes	ChipsRoastPots	Chips, roast potatoes	FALSE	TRUE
Fried potatoes	Chips	Chips, retail, fried in vegetable oil	TRUE	FALSE
Fried potatoes	PotatoSalad	Potato salad	TRUE	TRUE
Fried potatoes	RoastPotatoes	Old potatoes, roast in blended oil	TRUE	FALSE
Fruit juice	Smoothies	Smoothies (cup)	FALSE	TRUE
Fruit juice	PureFruitJuice	Pure fruit juice (100%) e.g. orange, apple juice (cup)	TRUE	TRUE
Grapes	Grapes	Grapes (handful)	TRUE	TRUE
Green leafy vegetables	GreenSalad	Green salad, lettuce, cucumber, celery	TRUE	TRUE
Green leafy vegetables	Spinach	Spinach	TRUE	TRUE
Green leafy vegetables	Watercress	Watercress	TRUE	TRUE
Herbal tea	FruitTea	Fruit tea (cup)	FALSE	TRUE
Herbal tea	GreenTea	Green tea (cup)	FALSE	TRUE
High fat milk	ChannellslandMilk	Channel Islands milk	TRUE	TRUE
High fat milk	EvaporatedMilk	Evaporated milk	TRUE	TRUE
High fat milk	FullMilk	Full cream milk	TRUE	TRUE
High fat milk	CoffeeWhitener	Coffee whitener, e.g. Coffee- mate (teaspoon)	TRUE	TRUE
High fat salad dressing	FrenchDressing	French dressing (tablespoon)	TRUE	TRUE
High fat salad	Мауо	Full fat salad cream, mayonnaise	TRUE	TRUE
High fat salad	Otherdressing	Other salad dressing (tablespoon)	TRUE	TRUE
High fibre breakfast cereals	HighFibreCereal	High Fibre cereals e.g. Branflakes, All Bran, Fruit and Fibre	FALSE	TRUE
High fibre breakfast cereals	AllBran	All Bran	TRUE	FALSE
High fibre breakfast cereals	Branflakes	Branflakes	TRUE	FALSE
High fibre breakfast cereals	Cheerios	Cheerios	TRUE	FALSE
High fibre breakfast cereals	FruitnFibre	Fruit n' fibre	TRUE	FALSE
High fibre breakfast cereals	Grapenuts	Grape nuts	TRUE	FALSE
High fibre breakfast cereals	Muesli	Muesli	TRUE	TRUE
High fibre breakfast cereals	OatBasedCereal	Oat based	TRUE	FALSE

Food group	FEO Variable Name	EEO Description	Included	Included
High fibre	OtherCereal	Cereal other	TRUE	FALSE
breakfast cereals	ounoroonour			171202
High fibre breakfast cereals	ShreddedWheat	Shredded wheat	TRUE	FALSE
High fibre breakfast cereals	Shreddies	Shreddies	TRUE	FALSE
High fibre breakfast cereals	Start	Start	TRUE	FALSE
High fibre breakfast cereals	SultanaBran	Sultana Bran	TRUE	FALSE
High fibre	WeetabixType	Weetabix type	TRUE	FALSE
High fibre	WheatFlakes	Wheat flakes	TRUE	FALSE
High sugar drinks	FruitSquash	Fruit squash or cordial (cup)	TRUE	TRUE
High sugar drinks	FizzySoftDrinks	Fizzy soft drinks, e.g. Coca Cola,	TRUE	TRUE
Ice cream	IceCream	Ice cream, choc ices	TRUE	TRUE
Lasagne	Lasagne	Lasagne, moussaka	TRUE	TRUE
Legumes	Beansprouts	Beansprouts	TRUE	TRUE
Legumes	DriedLentils	Pulses e.g. lentils, beans, peas	TRUE	TRUE
Legumes	GreenBeans	Green beans, broad beans, runner beans	TRUE	TRUE
Legumes	Peas	Peas	TRUE	TRUE
Low fat milk	DriedMilk	Dried milk	TRUE	TRUE
Low fat milk	SemiSkimmedMilk	Semi-skimmed milk	TRUE	TRUE
Low fat milk	SkimmedMilk	Skimmed milk	TRUE	TRUE
Low fat salad dressing	Lowcalsaladcream	Low calorie, low fat salad cream (tablespoon)	TRUE	TRUE
Low fat spread	LowFatSpread	Low fat spread, e.g. Outline, Gold (teaspoon)	TRUE	TRUE
Low fat spread	VLowFatSpread	Very low fat spread (teaspoon) e.g. Diet Flora	TRUE	TRUE
Low fibre breakfast cereals	BreakfastCereal	Breakfast cereal e.g. Cornflakes, Rice Krispies	FALSE	TRUE
Low fibre breakfast cereals	CocoPops	Coco pops	TRUE	FALSE
Low fibre breakfast cereals	Cornflakes	Corn Flakes	TRUE	FALSE
Low fibre	CrunchynutCornflake	Crunchynut cornflakes	TRUE	FALSE
Low fibre	Frosties	Sugar topped cereals e.g. Frosties	TRUE	TRUE
Low fibre breakfast cereals	JustRightType	Just right type	TRUE	FALSE
Low fibre	RiceKrispies	Rice Krispies	TRUE	FALSE
Low fibre	SpecialK	Special K	TRUE	FALSE
Low fibre	SugarPuffType	Sugar puffs	TRUE	FALSE
Low sugar soda	Dietfizzy	Low calorie or diet fizzy soft drinks (cup)	TRUE	TRUE
Malt drinks	Horlicks	Horlicks, Ovaltine (cup)	TRUE	TRUE
Margarine	CholLowerSpread	Cholesterol lowering fat spreads	FALSE	TRUE
Margarine	OliveOilSpread	Olive oil spread (teaspoon)	FALSE	TRUE
Margarine	BlockMarg	Block margarine, e.g. Stork, Krona (teaspoon)	TRUE	TRUE

Food group	FFQ Variable Name	FEQ Description	Included	Included
Margarine	OtherSoftMarg	Other soft margarine, dairy	TRUE	TRUE
	3	spreads, e.g. Blue Band, Clover (teaspoon)		
Marrow	Marrow	Marrow, courgettes	TRUE	TRUE
Meat	MFBeef	Beef: roast, steak, mince, stew or casserole	TRUE	TRUE
Meat	MFLamb	Lamb: roast, chops or stew	TRUE	TRUE
Meat	MFPork	Pork: roast, chops or stew	TRUE	TRUE
Melon	Melon	Melon (1 slice)	TRUE	TRUE
Mushrooms	Mushrooms	Mushrooms	TRUE	TRUE
Nuts	NutsSalted	Salted nuts e.g. peanuts, cashews (handful)	FALSE	TRUE
Nuts	NutsUnsalted	Unsalted nuts, e.g. brazil, walnuts (handful)	FALSE	TRUE
Nuts	Seeds	Seeds e.g. Sunflower, pumpkin (tablespoon)	FALSE	TRUE
Nuts	Nuts	Mixed nuts	TRUE	FALSE
Nuts	PeanutButter	Peanut butter (teaspoon)	TRUE	TRUE
Oily fish	MFOilyfish	Oily fish, fresh or canned, e.g. tuna, mackerel, kippers, salmon, sardines, herring	TRUE	TRUE
Other fish/seafood	MFFishroe	Fish roe, taramasalata	TRUE	TRUE
Other fish/seafood	MFShellfish	Shellfish, e.g. crab, prawns, mussels	TRUE	TRUE
Other fish/seafood	MFWhitefish	Other white fish, fresh or frozen, e.g. cod, plaice, sole, haddock, halibut	TRUE	TRUE
Peaches	Peaches	Peaches, plums, apricots (1 fruit)	TRUE	TRUE
Pizza	Pizza	Pizza (one slice)	TRUE	TRUE
Pizza	Quiche	Quiche (slice)	TRUE	TRUE
Polyunsaturated margarine	PufaMarg	Polyunsaturated margarine, e.g. Flora, sunflower (teaspoon)	TRUE	TRUE
Porridge	Porridge	Porridge, Readybreak, oats	TRUE	TRUE
Poultry	MFPoultry	Chicken or other poultry e.g. turkey	TRUE	TRUE
Processed fruit	DriedFruit	Dried fruit, e.g. raisins, prunes (heaped tablespoon)	TRUE	TRUE
Processed fruit	TinnedFruit	Tinned fruit (handful)	TRUE	TRUE
Processed meats	MFBacon	Bacon or gammon	TRUE	TRUE
Processed meats	MFCornedBeef	Corned Beef, Spam, luncheon meats	TRUE	TRUE
Processed meats	MFHam	Ham, cured meats & chorizo	TRUE	TRUE
Processed meats	MFLiver	Liver, liver pate, liver sausage	TRUE	TRUE
Processed meats	MFSausages	Sausages	TRUE	TRUE
Root vegetables	Beetroot	Beetroot	TRUE	TRUE
Root vegetables	Parsnips	Parsnips, turnips, swedes	TRUE	TRUE
Root vegetables	Carrots	Carrots	TRUE	TRUE
Savoury pies	MFPies	Savoury pies, e.g. meat pie, pork pie, pasties, steak & kidney pie, sausage rolls	TRUE	TRUE
Savoury snacks	CreamCrackers	Cream crackers, savoury biscuits	TRUE	TRUE
Savoury snacks	Crisps	Crisps or other packet snacks,	TRUE	TRUE

Food group	FEQ Variable Name	FFQ Description	Included	Included
Seasonings	Marmite	Marmite, Bovril (teaspoon)	TRUE	TRUE
Seasonings	PicklesChutney	Pickles, chutney (tablespoon)	TRUE	TRUE
Seasonings	Sauces	Sauces, e.g. white sauce, cheese sauce, gravy (tablespoon)	TRUE	TRUE
Seasonings	TomatoKetchup	Tomato ketchup (tablespoon)	TRUE	TRUE
Soup	MeatSoup	Meat soups (bowl) (to include meat and vegetable soups)	TRUE	TRUE
Soup	VegSoup	Vegetable soups (bowl)	TRUE	TRUE
Soy foods	Tofu	Meat substitutes e.g. tofu, soyameat, textured vegetable protein, vegeburger	TRUE	TRUE
Soy/other milk	GoatsMilk	Goats' milk	TRUE	TRUE
Soy/other milk	RiceMilk	Rice milk	TRUE	TRUE
Soy/other milk	SoyaMilk	Soya milk	TRUE	TRUE
Spirits/liquor	Liqueurs	Port, sherry, vermouth, liqueurs (pub measure)	TRUE	TRUE
Spirits/liquor	Spirits	Špirits, e.g. gin, brandy, whisky, vodka (pub measure)	TRUE	TRUE
Sweet baked products	BiscuitsRedFat	Reduced fat biscuits e.g. Go Ahead, Highlights (one small packet or one small bar/biscuit)	FALSE	TRUE
Sweet baked products	Biscuitschoc	Sweet biscuits, chocolate, e.g. digestive (one)	TRUE	TRUE
Sweet baked products	Biscuitsplain	Sweet biscuits, plain, e.g. Nice, ginger (one)	TRUE	TRUE
Sweet baked products	Bunshome	Buns, pastries e.g. scones, flapjacks, croissants, doughnuts, home baked	TRUE	TRUE
Sweet baked products	Cakeshome	Cakes e.g. fruit, sponge, home baked	TRUE	TRUE
Sweet baked products	Cakesready	Cakes e.g. fruit, sponge, ready made	TRUE	TRUE
Sweet baked products	Fruitpiehome	Fruit pies, tarts, crumbles, home baked	TRUE	TRUE
Sweet baked products	Fruitpiesready	Fruit pies, tarts, crumbles, ready made	TRUE	TRUE
Sweet baked products	MilkPuddings	Milk puddings e.g. rice, custard, trifle	TRUE	TRUE
Sweet baked products	SpongePudhome	Sponge puddings, home baked	TRUE	TRUE
Sweet baked products	SpongePudready	Sponge puddings, ready made	TRUE	TRUE
Sweet peppers	SweetPeppers	Sweet peppers	TRUE	TRUE
Sweetcorn	Sweetcorn	Sweetcorn	TRUE	TRUE
Tomatoes	Tomatoes	Tomatoes	TRUE	TRUE
White/brown bread, refined grains	NaanPoppdmTort	Naan, poppadoms, flour tortillas	FALSE	TRUE
White/brown bread, refined grains	BrownBread	Brown bread/rolls	TRUE	TRUE
White/brown bread, refined grains	WhiteBread	White bread/rolls	TRUE	TRUE
White/brown bread, refined grains	Whitepasta	White or green pasta, e.g. spaghetti, macaroni, noodles	TRUE	TRUE

			Included	Included
Food group	FFQ Variable Name	FFQ Description	FFQ1	FFQ2
White/brown bread, refined grains	WhiteRice	White rice	TRUE	TRUE
Wholemeal bread/grains	BrownRice	Brown rice	TRUE	TRUE
Wholemeal bread/grains	WholemealBread	Wholemeal & granary bread/rolls	TRUE	TRUE
Wholemeal bread/grains	WholemealPasta	Wholemeal pasta	TRUE	TRUE
Wine	RedWine	Red wine (small glass)	FALSE	TRUE
Wine	WhiteWine	White wine (small glass)	FALSE	TRUE
Wine	Wine	White wine, red wine	TRUE	FALSE
Wine	Wine	White wine, red wine	TRUE	FALSE
Yoghurt	DairyFFYog	Full fat or Greek yoghurt (small pot)	TRUE	TRUE
Yoghurt	DairyLFYog	Low fat yoghurt, fromage frais (small pot)	TRUE	TRUE

Table 2. Food frequency questionnaire variables included in each food group anddescriptions

Notes: Details of each food frequency questionnaire (FFQ) variables included in each food group analysis are shown. FFQs completed before 2007 (FFQ1) included slightly different variables than FFQs completed during 2007 (FFQ2), therefore the right hand columns indicate if a particular variable was included in the final (TRUE) or not (FALSE).



al., 2007))	-		T		
Food	Fruit and		I raditional English	Dieting	Low Meat
Baked beans	01 (04,.01)	07 (18,.10)	.17 (–.01,.25)	.10 (19,.36)	.31 (16,.37)
Beefburgers	–.14 (–.16,–.13)	.13 (–.11,.23)	.18 (–.02,.25)	.15 (–.02,.23)	.08 (–.18,.19)
Beer	03 (05,01)	.22 (.07,.26)	.00 (–.18,.17)	.09 (03,.16)	.04 (–.12,.14)
Berries	.16 (.14,.19)	05 (11,.04)	05 (11,.04)	.00 (–.10,.09)	05 (13,.06)
Butter	07 (09,05)	.11 (.00,.19)	03 (16,.13)	35 (40,03)	20 (40,.37)
Citrus fruit	.19 (.16,.21)	11 (16,01)	01 (12,.09)	.08 (–.11,.16)	07 (15,.06)
Coffee	01 (04,.01)	.17 (.04,.21)	01 (17,.14)	.13 (–.20,.23)	16 (25,.06)
Cooked potatoes	.03 (.00,.05)	19 (27,.00)	.19 (–.02,.28)	08 (17,.08)	02 (15,.15)
Crisp bread	.11 (.09,.14)	06 (11,.02)	03 (10,.05)	.03 (–.07,.10)	.02 (–.08,.11)
Dairy products; high fat	07 (09,06)	.14 (.00,.20)	13 (21,.02)	12 (20,.02)	07 (18,.16)
Dairy products;	.15 (.13,.17)	29 (33,06)	05 (24,.21)	.25 (09,.29)	02 (27,.22)
Drinks; other	.04 (.02,.06)	07 (14,.07)	–.10 (–.17,.02)	.17 (–.01,.23)	.11 (–.19,.23)
Eggs	.00 (02,.02)	.04 (08,.12)	.11 (.01,.17)	05 (14,.07)	–.06 (–.16,.11)
Fried fish	–.11 (–.13,–.09)	–.02 (–.18,.13)	.21 (.08,.24)	–.01 (–.11,.10)	.03 (–.07,.13)
Fried potatoes	–.21 (–.22,–.19)	.09 (–.17,.23)	.22 (.04,.27)	.01 (12,.17)	.12 (–.05,.19)
Fruit juice	.09 (.07,.12)	.06 (03,.12)	07 (14,.03)	.08 (08,.13)	04 (13,.07)
High fibre breakfast cereals	.06 (.04,.09)	23 (29,.04)	14 (26,.12)	.13 (–.17,.23)	13 (27,.08)
Lasagne	.01 (–.01,.03)	.13 (.02,.18)	.00 (–.14,.13)	.25 (–.00,.31)	.13 (–.27,.29)
Legumes	.19 (.16,.21)	.03 (24,.23)	.29 (.10,.33)	16 (27,.18)	.13 (–.09,.28)
Low fibre breakfast cereals	07 (09,05)	06 (11,.01)	.05 (–.04,.11)	–.05 (–.12,.10)	.07 (03,.14)
Low fat spread	.02 (01,.04)	10 (14,01)	.02 (09,.12)	.15 (–.01,.22)	.11 (–.18,.22)
Margarine	–.09 (–.11,–.07)	01 (08,.07)	.07 (–.01,.12)	.00 (–.12,.16)	.13 (–.04,.20)
Meat	10 (12,08)	02 (27,.21)	.34 (.11,.41)	.05 (–.35,.31)	32 (39,.02)
Nuts	.06 (.02,.09)	0.14 (02,.24)	–.15 (–.25,.01)	12 (20,.11)	.07 (–.09,.18)
Oily fish	.15 (.12,.19)	0.01 (04,.07)	01 (09,.07)	.02 (–.16,.13)	12 (17,.02)
Other fish and seafood	.18 (.16,.21)	0.07 (03,.13)	.07 (–.06,.18)	04 (25,.19)	20 (26,.07)
Other fruit	.27 (.25,.29)	21 (26,.00)	07 (21,.13)	.09 (06,.14)	–.01 (–.13,.10)
Pizza	06 (08,04)	0.15 (.02,.22)	10 (21,.07)	.18 (–.09,.31)	.22 (20,.32)
Polyunsaturated margarine	02 (04,.00)	08 (14,.03)	06 (15,.06)	.07 (–.10,.21)	.18 (–.10,.25)
Porridge	.08 (.06,.10)	–.13 (–.17,–.03)	.05 (–.08,.15)	14 (20,.05)	.00 (–.15,.17)
Poultry	.05 (.03,.07)	02 (13,.09)	.14 (.01,.25)	.17 (–.30,.32)	25 (33,.04)
Processed meats	10 (12,08)	.02 (25,.24)	.32 (.10,.38)	.13 (–.23,.28)	16 (27,.06)
Salad dressing high fat	.12 (.09,.14)	.30 (.08,.34)	–.11 (–.31,.13)	02 (10,.08)	03 (10,.07)
Salad dressing low fat	.14 (.10,.17)	02 (13,.11)	.10 (–.03,.19)	.18 (–.03,.27)	.13 (–.20,.25)
Savoury pies	–.19 (–.21,–.17)	.03 (20,.19)	.24 (.09,.27)	.00 (10,.09)	01 (09,.09)
Savoury snacks	10 (12,08)	.12 (.03,.17)	03 (12,.08)	.04 (07,.13)	.09 (07,.15)
Seasonings	.02 (.00,.05)	.05 (–.05,.11)	.07 (04,.14)	.02 (–.15,.23)	.18 (–.06,.25)
Soda; high sugar	08 (10,06)	.09 (–.01,.15)	.04 (07,.12)	.08 (05,.19)	.11 (–.12,.20)
Soda; low sugar	.02 (01,.04)	.06 (–.05,.14)	.04 (–.10,.18)	.33 (–.05,.36)	.08 (34,.33)
Soup	.12 (.10,.14)	03 (10,.03)	.03 (–.05,.11)	.05 (17,.18)	13 (20,.02)
Soy and other milk	.05 (.03,.08)	.00 (06,.08)	03 (11,.05)	15 (22,.12)	.08 (–.10,.21)
Soy foods	.11 (.07,.14)	.06 (06,.17)	09 (24,.05)	15 (37,.40)	.38 (.03,.43)
Spirits and liquor	.00 (02,.02)	.20 (.06,.23)	.01 (–.17,.16)	.10 (–.18,.20)	13 (21,.04)

Table 3. Factor loadings for the five principal components (modified from (Te	ucher et
al. 2007))	

uii, 2001 //					
	Fruit and		Traditional		
Food	Vegetable	High Alcohol	English	Dieting	Low Meat
Sweet baked	–.16 (–.17,–.14)	22 (26,.00)	–.10 (–.25,.14)	22 (27,.10)	.00 (20,.24)
Sweets and sweet condiments	–.17 (–.19,–.15)	02 (08,.05)	01 (13,.09)	26 (29,.17)	.09 (–.19,.28)
Теа	03 (05,01)	23 (28,05)	.08 (–.13,.26)	24 (29,.12)	.02 (22,.27)
Vegetables; allium	.23 (.21,.25)	.27 (.04,.30)	.06 (–.18,.23)	–.13 (–.18,.07)	.02 (–.13,.18)
Vegetables; cruciferous	.21 (.17,.24)	03 (32,.26)	.36 (.14,.39)	08 (20,.15)	.07 (08,.19)
Vegetables; green leafy	.30 (.28,.32)	.12 (–.04,.17)	.07 (–.06,.15)	06 (12,.10)	.07 (04,.13)
Vegetables; other	.32 (.30,.33)	.18 (–.09,.26)	.15 (–.05,.25)	–.13 (–.20,.15)	.12 (–.07,.19)
Vegetables; yellow	.30 (.28,.32)	–.03 (–.17,.13)	.18 (.06,.23)	–.09 (–.15,.11)	.06 (–.07,.15)
White and brown bread, refined grains	–.11 (–.13,–.09)	.04 (02,.09)	03 (10,.05)	02 (14,.17)	.15 (–.04,.22)
Wholemeal bread and grains	.15 (.12,.17)	–.11 (–.22,.12)	18 (23,.00)	01 (10,.08)	01 (10,.08)
Wine	.08 (.06,.10)	.33 (.07,.37)	–.14 (–.35,.14)	.03 (–.25,.21)	22 (27,.02)

Table 3. Factor	[.] loadings for the	five principal	components	(modified from	(Teucher et
al., 2007))	-		-	-	-

Fruit & n	uts	Vegetable	9	Mea	t	Fish		Dairy		Cereal	ereal ¹ Legu		me
ltem	g	Item	g	ltem	g	ltem	g	Item	g	ltem	g	ltem	g
								Channel				Baked	
Apples	100	Avocado	75	Bacon	25	Shellfish	60	Islands milk Cottage	585	Brown bread	36	beans	135
Bananas	100	Beansprouts	20	Beef	128	Oilyfish	96	cheese	55	Brown rice Cream	180	Lentils	70
Dried fruit	30	Beetroot	40	Burgers Corned	78	Fried fish	180	Dairy cheese	40	crackers	7		
Tinned fruit	100	Broccoli	85	beef	30	Fish roe	45	Dairy deserts	125	Crispbread	10		
Grapefruit	80	Cabbage	95	Ham	23	Fish fingers	50	Low fat yogurt	125	Frosties	40		
Grapes	80	Brussel sprouts	90	Lamb	94	Whitefish	120	Double cream	30	Lasagne	420		
Melon	200	Carrots	60	Liver	40			Single cream	15	White pasta	230		
Oranges	120	Cauliflower	90	Meat pies	124			Dried milk	585	Porridge	160		
Peaches	70	Coleslaw	45	Pork	128			Eggs	50	Muesli Wholemeal	60		
Pears	170	Garlic	5	Poultry	125			Full milk	585	pasta	230		
Strawberries	100	Green beans	90	Sausages	40			Goats milk	585	White bread	36		
Mixed nuts ² Nuts	40	Green salad	30					Rice milk Semi-	585	White rice Wholemeal	180		
unsalted ²	40	Leeks	75					skimmed milk	585	bread	36		
Nuts salted ²	40	Marrow	90					Skimmed milk	585				
		Mushrooms	56					Soya milk Evaporated	585				
		Onions	60					milk	585				
		Peas	70										
		Spinach	90										
		Sweetcorn	85										
		Sweet peppers	80										
		Tomatoes	85										
		Watercress	20										
		Tofu	50										
		Parsnips	65										

Table 4. Items and gram amounts per frequency category for the Mediterranean diet score calculation

Notes: ¹The cereal category for FFQ 1 included multiple cereal types which were omitted from FFQ 2, therefore only items which were included in both questionnaires were quantified.

²'Mixed nuts' was quantified on only FFQ 1, 'nuts unsalted' and 'nuts salted' were quantified on FFQ 2. For this analysis, frequencies for these items were combined to make one category.

			Super	Discov	very	Discore	dant	Meta-ana	lysis	Whole sample
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Ρ	Beta(SE)	Р	R ²
Allium	X-11858			0.040(0.007)	3.65E-08	0.036(0.015)	1.69E-02	0.039(0.007)	1.42E-09	0.030
Allium vegetables	tryptophan betaine	Tryptophan metabolism	Amino acid	0.041(0.007)	3.82E-08	0.023(0.010)	1.57E-02	0.034(0.006)	4.73E-09	0.025
Allium	X-02269			0.024(0.005)	2.82E-07	0.019(0.010)	4.75E-02	0.023(0.004)	3.55E-08	0.010
Apples/pears	threitol	Nucleotide sugars, pentose metabolism	Carbo- hydrate	0.034(0.004)	2.50E-17	0.030(0.007)	4.05E-05	0.033(0.003)	1.69E-21	0.031
Apples/pears	X-11315			0.035(0.004)	9.14E-15	0.036(0.008)	6.10E-06	0.035(0.004)	9.63E-20	0.029
Apples/pears	X-11372			-0.024(0.004)	1.97E-09	-0.021(0.007)	3.40E-03	-0.023(0.003)	2.02E-11	0.014
Apples/pears	indolepropionate	Tryptophan metabolism	Amino acid	0.027(0.005)	4.50E-07	0.024(0.008)	1.71E-03	0.026(0.004)	2.39E-09	0.016
Apples/pears	3-phenylpropionate (hydrocinnamate)	Phenylalanine & tyrosine metabolism	Amino acid	0.025(0.005)	3.76E-07	0.023(0.009)	1.21E-02	0.024(0.004)	1.24E-08	0.015
Apples/pears	X-09789			0.021(0.004)	2.73E-07	0.016(0.007)	2.58E-02	0.020(0.004)	2.22E-08	0.010
Avocado	X-11469			0.204(0.027)	3.59E-14	0.100(0.027)	3.31E-04	0.153(0.019)	9.36E-16	0.023
Avocado	X-02269			0.193(0.028)	3.88E-12	0.107(0.026)	6.44E-05	0.147(0.019)	6.33E-15	0.021
Avocado	3-carboxy-4-methyl- 5-propyl-2- furanpropanoate (CMPF)	Fatty acid, dicarboxylate	Lipid	0.185(0.027)	8.83E-12	0.116(0.029)	1.26E-04	0.153(0.020)	1.03E-14	0.022
Avocado	1- docosahexaenoylgl ycerophosphocholi ne*	Lysolipid	Lipid	0.125(0.024)	2.54E-07	0.094(0.028)	1.06E-03	0.112(0.018)	1.03E-09	0.009
Avocado	eicosapentaenoate (FPA: 20:5n3)	Essential fatty acid	Lipid	0.163(0.026)	3.82E-10	0.056(0.025)	2.56E-02	0.108(0.018)	2.01E-09	0.013
Bananas	indolepropionate	Tryptophan metabolism	Amino acid	0.038(0.006)	5.61E-11	0.022(0.010)	3.37E-02	0.034(0.005)	1.05E-11	0.016
Beef burgers	trans-4- hydroxyproline	Urea cycle; arginine-, proline-, metabolism	Amino acid	0.428(0.085)	5.90E-07	0.257(0.104)	1.48E-02	0.360(0.066)	5.34E-08	0.009
Berries	X-11315			0.124(0.018)	2.28E-12	0.072(0.021)	8.29E-04	0.103(0.013)	2.70E-14	0.021
Black tea	X-14473			-0.025(0.001)	8.51E-60	-0.020(0.003)	5.16E-09	-0.024(0.001)	1.36E-72	0.110
Black tea	quinate	Food component, Plant	Xenobiotics	-0.019(0.002)	9.74E-34	-0.019(0.003)	1.61E-09	-0.019(0.001)	5.58E-44	0.066
Black tea	X-12816			-0.019(0.002)	4.68E-23	-0.021(0.004)	2.59E-07	-0.020(0.002)	3.57E-30	0.062

			Super-	Discovery		Discor	dant	Meta-ana	alysis	Whole sample
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Ρ	Beta(SE)	Р	R ²
Black tea	X-14374			-0.013(0.001)	8.48E-21	-0.011(0.003)	2.12E-03	-0.013(0.001)	2.50E-23	0.030
Black tea	X-120393- hydroxypyridine sulfate	Chemical	Xenobiotics	-0.014(0.002)	2.23E-16	-0.013(0.004)	4.61E-04	-0.014(0.002)	1.60E-19	0.035
Black tea	X-05426			-0.010(0.002)	8.54E-12	-0.008(0.003)	1.66E-02	-0.010(0.001)	3.62E-13	0.019
Black tea	cyclo(leu-pro)	Dipeptide	Peptide	-0.012(0.002)	5.64E-11	-0.010(0.004)	2.26E-02	-0.012(0.002)	2.89E-12	0.026
Black tea	X-12230			-0.009(0.002)	3.07E-08	-0.010(0.003)	4.67E-03	-0.009(0.001)	3.98E-10	0.015
Black tea	X-137413-methyl catechol sulfate 1	Benzoate metabolism	Xenobiotics	-0.010(0.002)	6.29E-08	-0.011(0.004)	4.92E-03	-0.010(0.002)	8.73E-10	0.017
Butter	X-13431 nonanoylcarnitine*	Carnitine metabolism	Lipid	0.028(0.004)	9.12E-15	0.020(0.006)	1.97E-03	0.026(0.003)	6.48E-17	0.026
Butter	X-02249			0.027(0.003)	6.97E-15	0.019(0.007)	1.06E-02	0.026(0.003)	2.31E-16	0.023
Butter	X-10510			0.023(0.003)	1.85E-11	0.019(0.005)	1.50E-04	0.022(0.003)	8.28E-15	0.019
Butter	15-methylpalmitate (isobar with 2- methylpalmitate)	Fatty acid, branched	Lipid	0.027(0.004)	6.27E-11	0.019(0.008)	1.38E-02	0.025(0.004)	3.09E-12	0.026
Butter	10-undecenoate (11:1n1)	Medium chain fatty acid	Lipid	0.023(0.004)	5.99E-10	0.014(0.005)	4.11E-03	0.020(0.003)	2.35E-11	0.018
Butter	myristate (14:0)	Long chain fatty acid	Lipid	0.019(0.004)	5.57E-08	0.019(0.006)	4.34E-03	0.019(0.003)	6.68E-10	0.014
Butter	X-08402			0.018(0.003)	1.85E-08	0.013(0.005)	1.10E-02	0.017(0.003)	7.17E-10	0.012
Butter	10-nonadecenoate (19:1n9)	Long chain fatty acid	Lipid	0.017(0.004)	9.50E-07	0.019(0.006)	8.91E-04	0.018(0.003)	2.43E-09	0.013
Butter	pentadecanoate (15:0)	Long chain fatty acid	Lipid	0.019(0.004)	8.48E-08	0.015(0.006)	1.03E-02	0.018(0.003)	2.77E-09	0.013
Chocolate	theobromine	Xanthine metabolism	Xenobiotics	0.024(0.005)	3.43E-07	0.023(0.005)	1.83E-05	0.024(0.003)	1.34E-11	0.020
Chocolate	7-methylxanthine	Xanthine metabolism	Xenobiotics	0.029(0.005)	2.48E-08	0.012(0.006)	3.31E-02	0.022(0.004)	2.00E-08	0.020
Citrus fruit	stachydrine	Food component, Plant	Xenobiotics	0.058(0.006)	6.43E-24	0.036(0.008)	4.05E-05	0.051(0.005)	2.07E-27	0.051
Citrus fruit	glycerate	Glycolysis, gluconeogenesis, pyruvate metabolism	Carbohydrat e	0.027(0.005)	1.15E-07	0.031(0.008)	1.39E-04	0.028(0.004)	4.57E-11	0.014
Citrus fruit	X-11315			0.029(0.005)	4.72E-08	0.021(0.007)	3.30E-03	0.026(0.004)	6.50E-10	0.014
Coffee	X-14473			0.041(0.001)	6.89E-139	0.027(0.003)	1.38E-19	0.038(0.001)	6.12E-187	0.246
Coffee	quinate	Food component, Plant	Xenobiotics	0.036(0.001)	6.60E-106	0.024(0.003)	6.84E-15	0.033(0.001)	4.31E-136	0.190

			C	Discov	very	Discore	dant	Meta-ana	lysis	Whole sample
Variable	Metabolite name	Pathway	Super- pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	R ²
Coffee	X-14374			0.025(0.002)	1.61E-55	0.019(0.003)	1.78E-10	0.023(0.001)	1.87E-68	0.083
Coffee	X-120393- hydroxypyridine sulfate	Chemical	Xenobiotics	0.026(0.002)	6.89E-52	0.020(0.003)	1.72E-09	0.025(0.001)	1.64E-64	0.108
Coffee	X-12816			0.032(0.002)	1.07E-51	0.021(0.004)	7.67E-07	0.030(0.002)	3.22E-61	0.133
Coffee	X-05426			0.019(0.002)	5.06E-34	0.012(0.003)	2.06E-04	0.018(0.001)	6.64E-38	0.058
Coffee	X-12230			0.019(0.002)	1.07E-27	0.015(0.003)	1.99E-06	0.018(0.002)	1.02E-33	0.060
Coffee	X-137413-methyl catechol sulfate 1	Benzoate metabolism	Xenobiotics	0.020(0.002)	3.43E-27	0.016(0.004)	1.05E-05	0.019(0.002)	1.25E-32	0.063
Coffee	catechol sulfate	Benzoate metabolism	Xenobiotics	0.015(0.002)	5.39E-22	0.011(0.003)	2.24E-04	0.014(0.001)	3.44E-25	0.033
Coffee	cyclo(leu-pro)	Dipeptide	Peptide	0.014(0.002)	9.25E-14	0.009(0.004)	1.58E-02	0.013(0.002)	5.29E-15	0.030
Coffee	X-12217O-methyl catechol sulfate	Benzoate metabolism	Xenobiotics	0.012(0.002)	5.43E-12	0.009(0.003)	8.66E-03	0.011(0.002)	1.45E-13	0.022
Coffee	1-methylxanthine	Xanthine metabolism	Xenobiotics	0.010(0.002)	2.82E-07	0.010(0.003)	4.20E-03	0.010(0.002)	3.31E-09	0.016
Confectionary /jam	X-11315			-0.011(0.002)	6.99E-11	-0.006(0.003)	2.77E-02	-0.010(0.001)	1.53E-11	0.017
Confectionary /jam	glycerate	Glycolysis, gluconeogenesis, pyruvate metabolism	Carbo- hydrate	-0.011(0.002)	4.86E-08	-0.008(0.003)	1.30E-02	-0.010(0.002)	2.02E-09	0.014
Confectionary /iam	pipecolate	Lysine metabolism	Amino acid	-0.009(0.002)	4.07E-07	-0.008(0.004)	3.81E-02	-0.008(0.002)	3.94E-08	0.008
Fried fish	X-11372			0.360(0.051)	2.88E-12	0.165(0.049)	8.66E-04	0.258(0.035)	2.81E-13	0.040
Fried fish	3-phenylpropionate (hydrocinnamate)	Phenylalanine & tyrosine metabolism	Amino acid	-0.167(0.032)	2.07E-07	-0.184(0.045)	7.67E-05	-0.172(0.026)	4.12E-11	0.012
Fried fish	X-11315			-0.174(0.035)	6.30E-07	-0.149(0.044)	7.69E-04	-0.164(0.027)	1.54E-09	0.013
Fried fish	scyllo-inositol	Inositol metabolism	Lipid	-0.170(0.031)	6.22E-08	-0.116(0.048)	1.77E-02	-0.154(0.026)	4.44E-09	0.011
Fruit juice	stachydrine	Food component, Plant	Xenobiotics	0.064(0.005)	2.87E-30	0.045(0.008)	1.06E-07	0.058(0.005)	3.26E-37	0.064
Green leafy vegetables	X-11469			0.034(0.005)	9.00E-11	0.025(0.006)	1.24E-04	0.030(0.004)	5.66E-14	0.020
Green leafy vegetables	X-02269			0.033(0.005)	9.43E-10	0.025(0.006)	6.56E-05	0.030(0.004)	2.39E-13	0.018
Green leafy vegetables	X-11315			0.038(0.006)	4.37E-10	0.025(0.007)	5.56E-04	0.033(0.005)	1.71E-12	0.023
Green leafy vegetables	X-11372			-0.028(0.004)	5.23E-10	-0.021(0.009)	1.58E-02	-0.026(0.004)	2.51E-11	0.012

			Discovery Super-		Discore	dant	Meta-ana	lysis	Whole sample	
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	R ²
Green leafy vegetables	3-carboxy-4-methyl- 5-propyl-2- furanpropanoate (CMPF)	Fatty acid, dicarboxylate	Lipid	0.028(0.006)	9.29E-07	0.021(0.006)	8.06E-04	0.025(0.004)	3.03E-09	0.013
Green leafy vegetables	1- docosahexaenoylgl ycerophosphocholi ne*	Lysolipid	Lipid	0.025(0.005)	9.36E-08	0.017(0.007)	1.87E-02	0.023(0.004)	7.11E-09	0.009
High fat salad dressing	eicosapentaenoate (EPA; 20:5n3)	Essential fatty acid	Lipid	0.048(0.010)	8.34E-07	0.035(0.015)	1.82E-02	0.044(0.008)	5.48E-08	0.010
High fibre breakfast cereals	X-09789			0.046(0.006)	4.90E-14	0.019(0.009)	4.34E-02	0.038(0.005)	7.28E-14	0.021
High fibre breakfast cereals	X-11469			0.032(0.005)	1.54E-10	0.028(0.008)	1.21E-03	0.031(0.004)	5.40E-13	0.013
High fibre breakfast cereals	X-02269			0.031(0.005)	7.68E-10	0.028(0.009)	1.65E-03	0.030(0.004)	3.63E-12	0.012
High fibre breakfast cereals	pyridoxate	Vitamin B6 metabolism	Cofactors and vitamins	0.040(0.006)	9.41E-11	0.020(0.010)	3.54E-02	0.035(0.005)	3.52E-11	0.019
High fibre breakfast cereals	X-11315			0.031(0.006)	8.40E-08	0.027(0.010)	1.12E-02	0.030(0.005)	2.75E-09	0.011
Low fat mik	X-21365 [trimethyl- N-aminovalerate]	Carnitine metabolism	Lipid	0.085(0.008)	3.30E-25	0.045(0.015)	2.55E-03	0.076(0.007)	9.36E-27	0.018
Low fat mik	X-12798			0.071(0.009)	4.88E-15	0.036(0.016)	2.12E-02	0.062(0.008)	1.24E-15	0.024
Meat	trans-4- hydroxyproline	Urea cycle; arginine-, proline-, metabolism	Amino acid	0.077(0.010)	7.70E-14	0.070(0.017)	6.28E-05	0.075(0.009)	1.08E-17	0.020
Meat	creatine	Creatine metabolism	Amino acid	0.070(0.011)	5.01E-11	0.041(0.019)	3.12E-02	0.063(0.009)	8.24E-12	0.016
Meat	pyroglutamine*	Glutamate metabolism	Amino acid	-0.066(0.011)	4.33E-09	-0.049(0.020)	1.46E-02	-0.062(0.010)	2.10E-10	0.016
Meat	X-09789			-0.052(0.011)	1.11E-06	-0.053(0.020)	7.66E-03	-0.052(0.009)	2.39E-08	0.009
Mushrooms	ergothioneine	Food component, Plant	Xenobiotics	0.189(0.023)	6.45E-16	0.165(0.033)	9.28E-07	0.181(0.019)	5.93E-22	0.107
Nuts	X-11315			0.060(0.006)	6.48E-21	0.042(0.009)	1.81E-05	0.054(0.005)	3.75E-25	0.037
Oily fish	docosahexaenoate (DHA: 22:6n3)	Essential fatty acid	Lipid	0.244(0.017)	1.72E-45	0.087(0.019)	9.66E-06	0.177(0.013)	2.09E-44	0.064
Oily fish	X-11469			0.214(0.017)	6.59E-34	0.117(0.022)	1.89E-07	0.176(0.013)	5.87E-39	0.062

			C	Discov	very	Discore	dant	Meta-ana	lysis	Whole sample
Variable	Metabolite name	Pathway	Super- pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	R ²
Oily fish	3-carboxy-4-methyl- 5-propyl-2- furanpropanoate (CMPF)	Fatty acid, dicarboxylate	Lipid	0.204(0.016)	3.52E-34	0.112(0.022)	9.22E-07	0.172(0.013)	8.57E-39	0.054
Oily fish	X-02269			0.208(0.017)	1.44E-32	0.122(0.022)	7.36E-08	0.175(0.013)	1.44E-38	0.058
Oily fish	eicosapentaenoate (EPA; 20:5n3)	Essential fatty acid	Lipid	0.212(0.017)	1.95E-33	0.085(0.024)	5.01E-04	0.169(0.014)	1.57E-33	0.052
Oily fish	1- docosahexaenoylgl ycerophosphocholi ne*	Lysolipid	Lipid	0.145(0.017)	7.40E-17	0.068(0.032)	3.36E-02	0.128(0.015)	3.33E-17	0.025
Oily fish	X-11315			0.138(0.018)	6.44E-15	0.054(0.022)	1.68E-02	0.106(0.014)	1.62E-14	0.020
Oily fish	1- oleoylglycerophosp hoethanolamine	Lysolipid	Lipid	-0.115(0.018)	2.65E-10	-0.074(0.026)	4.23E-03	-0.101(0.015)	6.89E-12	0.012
Oily fish	1- eicosatrienoylglycer ophosphocholine*	Lysolipid	Lipid	-0.099(0.017)	9.17E-09	-0.087(0.024)	4.03E-04	-0.095(0.014)	1.12E-11	0.011
Oily fish	1- arachidonoylglycer ophosphoethanola mine*	Lysolipid	Lipid	-0.106(0.018)	1.24E-08	-0.081(0.026)	1.97E-03	-0.097(0.015)	9.46E-11	0.013
Oily fish	docosapentaenoate (n3 DPA; 22:5n3)	Essential fatty acid	Lipid	0.111(0.017)	3.62E-11	0.038(0.019)	4.72E-02	0.080(0.013)	2.22E-10	0.014
Oily fish	1- linoleoylglyceropho sphoethanolamine*	Lysolipid	Lipid	-0.106(0.018)	5.45E-09	-0.055(0.025)	3.12E-02	-0.089(0.015)	1.60E-09	0.009
Oily fish	X-12627			0.093(0.016)	1.38E-08	0.044(0.019)	2.34E-02	0.073(0.012)	5.74E-09	0.009
Other fish/seafood	3-carboxy-4-methyl- 5-propyl-2- furanpropanoate (CMPF)	Fatty acid, dicarboxylate	Lipid	0.201(0.017)	1.16E-29	0.150(0.029)	8.90E-07	0.188(0.015)	6.64E-36	0.061
Other fish/seafood	docosahexaenoate (DHA; 22:6n3)	Essential fatty acid	Lipid	0.198(0.017)	5.83E-29	0.141(0.028)	9.10E-07	0.182(0.015)	6.10E-35	0.052
Other fish/seafood	X-11469			0.189(0.017)	5.51E-27	0.131(0.031)	3.07E-05	0.175(0.015)	2.99E-31	0.052
Other fish/seafood	X-02269			0.184(0.017)	7.33E-26	0.141(0.031)	1.21E-05	0.174(0.015)	9.25E-31	0.050
Other fish/seafood	eicosapentaenoate (EPA; 20:5n3)	Essential fatty acid	Lipid	0.156(0.017)	1.71E-19	0.148(0.033)	1.38E-05	0.154(0.015)	2.66E-24	0.036
Other fish/seafood	1- docosahexaenoylgl ycerophosphocholi ne*	Lysolipid	Lipid	0.125(0.015)	7.30E-16	0.071(0.029)	1.35E-02	0.113(0.014)	6.70E-17	0.020

			Sumar	Discov	very	Discor	dant	Meta-ana	alysis	Whole sample
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	R ²
Other fish/seafood	pyroglutamine*	Glutamate metabolism	Amino acid	-0.097(0.018)	3.81E-08	-0.102(0.025)	5.51E-05	-0.099(0.014)	5.22E-12	0.015
Other fish/seafood	docosapentaenoate (n3 DPA; 22:5n3)	Essential fatty acid	Lipid	0.086(0.015)	1.27E-08	0.102(0.030)	8.79E-04	0.089(0.013)	3.44E-11	0.010
Other fish/seafood	1- eicosatrienoylglycer ophosphocholine*	Lysolipid	Lipid	-0.082(0.016)	2.73E-07	-0.083(0.032)	1.14E-02	-0.082(0.014)	8.44E-09	0.009
Other fish/seafood	1,5-anhydroglucitol (1,5-AG)	Glycolysis, gluconeogenesis, pyruvate metabolism	Carbo- hydrate	-0.079(0.016)	6.12E-07	-0.087(0.034)	1.06E-02	-0.081(0.014)	1.80E-08	0.011
Other fish/seafood	X-126963,4- dihydroxyphenylace tate sulfate	Phenylalanine and Tyrosine Metabolism	Amino acid	-0.084(0.017)	6.20E-07	-0.078(0.036)	2.96E-02	-0.083(0.015)	4.73E-08	0.011
Peaches	X-11315			0.158(0.024)	8.45E-11	0.097(0.024)	6.51E-05	0.127(0.017)	5.92E-14	0.039
Porridge	X-09789			0.099(0.009)	9.10E-26	0.083(0.015)	1.01E-07	0.094(0.008)	4.96E-33	0.048
Porridge	X-122532- aminophenol sulfate	Chemical	Xenobiotics	0.053(0.010)	3.85E-08	0.043(0.019)	2.50E-02	0.051(0.009)	2.63E-09	0.016
Poultry	pyroglutamine*	Glutamate metabolism	Amino acid	-0.102(0.016)	1.87E-10	-0.070(0.022)	1.29E-03	-0.091(0.013)	1.36E-12	0.017
Poultry	creatine	Creatine metabolism	Amino acid	0.104(0.015)	2.57E-11	0.044(0.022)	4.56E-02	0.083(0.013)	3.17E-11	0.015
Processed meats	trans-4- hydroxyproline	Urea cycle; arginine-, proline-, metabolism	Amino acid	0.052(0.009)	4.07E-09	0.035(0.016)	3.14E-02	0.048(0.008)	4.69E-10	0.011
Processed meats	X-11381			0.052(0.009)	6.89E-09	0.039(0.017)	2.18E-02	0.049(0.008)	4.74E-10	0.011
Savoury pies	X-11315			-0.287(0.044)	5.59E-11	-0.116(0.054)	3.44E-02	-0.221(0.034)	9.08E-11	0.016
Savoury pies	3-phenylpropionate (hydrocinnamate)	Phenylalanine & tyrosine metabolism	Amino acid	-0.264(0.044)	2.06E-09	-0.151(0.068)	2.89E-02	-0.231(0.037)	3.72E-10	0.013
Savoury pies	X-11372			0.290(0.047)	1.14E-09	0.114(0.055)	4.02E-02	0.216(0.036)	2.04E-09	0.015
Savoury pies	ergothioneine	Food component, Plant	Xenobiotics	-0.320(0.060)	1.31E-07	-0.291(0.105)	6.63E-03	-0.313(0.052)	2.16E-09	0.017
Savoury snacks	X-11372			0.064(0.008)	8.31E-14	0.029(0.011)	9.73E-03	0.051(0.007)	3.88E-14	0.026
Soy foods	X-11381			-0.184(0.033)	1.95E-08	-0.063(0.025)	1.46E-02	-0.108(0.020)	5.80E-08	0.018
Soy/other milk	4- ethylphenylsulfate	Benzoate metabolism	Xenobiotics	0.299(0.051)	6.10E-09	0.195(0.043)	3.18E-05	0.239(0.033)	6.05E-13	0.061
Spirits/liquor	4-androsten- 3beta,17beta-diol	Sterol, Steroid	Lipid	0.044(0.008)	2.53E-08	0.028(0.013)	4.24E-02	0.040(0.007)	4.30E-09	0.021

			Discovery Super-		very	Discordant		Meta-ana	Whole sample	
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Ρ	Beta(SE)	Ρ	R ²
	disulfate 1*									
Sweet baked products	eicosapentaenoate (EPA; 20:5n3)	Essential fatty acid	Lipid	-0.016(0.002)	1.20E-10	-0.012(0.004)	2.30E-03	-0.014(0.002)	1.08E-12	0.016
Sweet baked products	docosahexaenoate (DHA; 22:6n3)	Essential fatty acid	Lipid	-0.016(0.002)	3.75E-11	-0.011(0.004)	7.33E-03	-0.015(0.002)	1.28E-12	0.016
Sweet baked products	scyllo-inositol	Inositol metabolism	Lipid	-0.018(0.003)	8.43E-09	-0.014(0.005)	5.30E-03	-0.017(0.003)	1.44E-10	0.019
Sweet baked	X-11799			-0.018(0.003)	1.39E-07	-0.015(0.006)	1.37E-02	-0.017(0.003)	4.88E-09	0.020
Sweet baked products	X-11315			-0.014(0.003)	3.66E-07	-0.009(0.003)	1.26E-02	-0.012(0.002)	2.90E-08	0.014
Tomatoes	glycerate	Glycolysis, gluconeogenesis, pyruvate metabolism	Carbo- hydrate	0.029(0.006)	7.50E-07	0.018(0.007)	1.02E-02	0.025(0.005)	4.72E-08	0.009
White/brown bread, refined grains	X-11315			-0.017(0.003)	1.00E-08	-0.019(0.005)	6.53E-05	-0.017(0.002)	1.84E-12	0.017
Wholemeal bread/grains	X-09789			0.017(0.003)	3.91E-07	0.011(0.005)	3.58E-02	0.015(0.003)	6.03E-08	0.010
Wine	scyllo-inositol	Inositol metabolism	Lipid	0.056(0.004)	3.04E-43	0.036(0.007)	3.42E-06	0.052(0.003)	1.47E-49	0.117
Wine	alpha- hydroxyisovalerate	Valine, leucine and isoleucine metabolism	Amino acid	0.047(0.004)	2.09E-36	0.038(0.006)	2.32E-08	0.045(0.003)	1.23E-45	0.072
Wine	X-11795			0.042(0.003)	5.04E-34	0.030(0.007)	9.51E-05	0.039(0.003)	1.74E-38	0.062
Wine	4-androsten- 3beta,17beta-diol disulfate 1*	Sterol, Steroid	Lipid	0.043(0.004)	1.09E-26	0.034(0.008)	3.31E-05	0.041(0.003)	1.50E-31	0.071
Wine	X-11799			0.035(0.004)	3.49E-22	0.028(0.006)	8.74E-06	0.033(0.003)	1.66E-27	0.052
Wine	X-10395			0.035(0.003)	9.31E-25	0.018(0.006)	3.51E-03	0.031(0.003)	3.45E-26	0.034
Wine	X-04495			0.032(0.003)	4.68E-20	0.028(0.006)	8.69E-06	0.031(0.003)	3.26E-25	0.035
Wine	5alpha-androstan- 3beta,17beta-diol disulfate	Sterol, Steroid	Lipid	0.038(0.004)	6.27E-21	0.030(0.008)	4.21E-04	0.036(0.004)	2.71E-24	0.059
Wine	pipecolate	Lysine metabolism	Amino acid	0.031(0.003)	1.57E-19	0.024(0.007)	4.94E-04	0.029(0.003)	1.51E-22	0.029
Wine	eicosapentaenoate (EPA; 20:5n3)	Essential fatty acid	Lipid	0.030(0.003)	1.66E-20	0.017(0.006)	4.40E-03	0.027(0.003)	6.63E-22	0.027
Wine	docosahexaenoate (DHA; 22:6n3)	Essential fatty acid	Lipid	0.029(0.003)	2.42E-17	0.014(0.005)	1.25E-02	0.025(0.003)	6.92E-18	0.022
Wine	myo-inositol	Inositol metabolism	Lipid	0.026(0.003)	7.58E-16	0.011(0.005)	4.93E-02	0.022(0.003)	1.36E-15	0.022

			Super-	Discovery		Discore	dant	Meta-ana	llysis	Whole sample
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	R ²
Wine	docosapentaenoate (n3 DPA; 22:5n3)	Essential fatty acid	Lipid	0.025(0.003)	5.38E-13	0.019(0.006)	3.46E-03	0.024(0.003)	5.28E-15	0.019
Wine	X-12627			0.024(0.003)	4.95E-14	0.014(0.006)	1.55E-02	0.022(0.003)	5.88E-15	0.017
Wine	X-01911			0.029(0.004)	7.11E-11	0.022(0.006)	1.27E-04	0.026(0.003)	2.55E-14	0.031
Wine	2-aminobutyrate	Butanoate metabolism	Amino acid	0.027(0.004)	2.83E-13	0.016(0.007)	2.04E-02	0.025(0.003)	2.73E-14	0.023
Wine	X-05907			0.024(0.003)	4.22E-12	0.017(0.005)	1.85E-03	0.022(0.003)	3.03E-14	0.019
Wine	X-10429			0.025(0.003)	1.88E-12	0.019(0.007)	7.10E-03	0.024(0.003)	3.62E-14	0.019
Wine	caprate (10:0)	Medium chain fatty acid	Lipid	0.021(0.003)	1.36E-11	0.023(0.007)	2.09E-03	0.022(0.003)	6.60E-14	0.016
Wine	piperine	Food component, Plant	Xenobiotics	0.028(0.004)	1.04E-10	0.022(0.006)	2.34E-04	0.026(0.003)	6.81E-14	0.026
Wine	X-12038			0.020(0.003)	2.27E-09	0.028(0.007)	1.20E-04	0.022(0.003)	9.20E-13	0.014
Wine	X-13215			0.023(0.004)	1.48E-09	0.023(0.006)	3.37E-04	0.023(0.003)	1.07E-12	0.016
Wine	X-11317			0.019(0.003)	3.11E-09	0.026(0.007)	1.84E-04	0.020(0.003)	1.88E-12	0.012
Wine	4-methyl-2- oxopentanoate	Valine, leucine and isoleucine metabolism	Amino acid	0.021(0.003)	1.36E-09	0.021(0.006)	1.07E-03	0.021(0.003)	3.44E-12	0.011
Wine	beta- hydroxyisovalerate	Valine, leucine and isoleucine metabolism	Amino acid	0.021(0.003)	9.19E-11	0.016(0.007)	1.64E-02	0.020(0.003)	4.44E-12	0.009
Wine	X-09026			0.020(0.003)	1.93E-09	0.019(0.006)	2.71E-03	0.020(0.003)	1.30E-11	0.011
Wine	caprylate (8:0)	Medium chain fatty acid	Lipid	0.019(0.003)	2.10E-09	0.020(0.007)	4.05E-03	0.019(0.003)	2.13E-11	0.012
Wine	benzoate	Benzoate metabolism	Xenobiotics	0.019(0.003)	4.37E-09	0.018(0.006)	2.84E-03	0.019(0.003)	3.20E-11	0.009
Wine	X-11550			0.018(0.003)	1.54E-08	0.020(0.006)	9.97E-04	0.018(0.003)	3.94E-11	0.011
Wine	stearidonate (18:4n3)	Long chain fatty acid	Lipid	0.019(0.003)	1.47E-08	0.019(0.007)	8.49E-03	0.019(0.003)	3.17E-10	0.012
Wine	2-hydroxybutyrate (AHB)	Cysteine, methionine, SAM, taurine metabolism	Amino acid	0.019(0.004)	4.61E-08	0.021(0.007)	3.45E-03	0.020(0.003)	4.15E-10	0.015
Wine	3-(4- hydroxyphenyl)lacta te	Phenylalanine & tyrosine metabolism	Amino acid	0.017(0.003)	3.11E-08	0.018(0.007)	1.37E-02	0.018(0.003)	1.08E-09	0.011
Wine	3-methyl-2- oxobutyrate	Valine, leucine and isoleucine metabolism	Amino acid	0.018(0.003)	2.72E-07	0.020(0.007)	3.55E-03	0.018(0.003)	2.61E-09	0.007

			Super-	Discov	ery	Discore	dant	Meta-ana	llysis	Whole sample
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Ρ	R ²
Wine	X-11452			0.023(0.004)	1.18E-07	0.017(0.006)	6.54E-03	0.021(0.003)	2.79E-09	0.015
Wine	epiandrosterone sulfate	Sterol, Steroid	Lipid	0.018(0.003)	2.21E-07	0.016(0.006)	1.05E-02	0.018(0.003)	6.29E-09	0.010
Wine	X-13496			0.017(0.003)	1.08E-07	0.015(0.006)	2.23E-02	0.017(0.003)	6.56E-09	0.009
Wine	arachidonate (20:4n6)	Long chain fatty acid	Lipid	0.017(0.004)	1.13E-06	0.018(0.006)	5.04E-03	0.017(0.003)	1.50E-08	0.009
Wine	theophylline	Xanthine metabolism	Xenobiotics	0.020(0.004)	2.47E-07	0.014(0.006)	2.41E-02	0.018(0.003)	1.99E-08	0.009
Wine	X-126441- docosahexaenoylgl ycerophosphoethan olamine	Lysolipid	Lipid	0.015(0.003)	8.33E-07	0.017(0.007)	2.16E-02	0.015(0.003)	4.82E-08	0.007

Notes: Table shows results of the linear regression analysis for the discovery population (excluding monozygotic twins discordant for each food group intake), the MZ discordant twin sample and the fixed effects meta-analysis of both groups. Only significant associations are shown which includes those associations passing the bonferroni cut-off in the discovery and fixed effects analyses ($1.08 \times 10^{-6} = 0.05/[77 food variables \times 601$ detected metabolites]) and passing the 5% level of significance in the discordant twin group. The R-squared was calculated for the relevant metabolite and food group in the study population. The Metabolon platform is a non-targeted platform which identified 456 metabolites in blood; data were available for 3559 twins.

			Supar	Discov	ery	Discor	dant	Meta-ana	alysis	Whole sample
Variable	Biocrates name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	R ²
Cream	lysoPhosphatidylcholine acyl C17:0	Glycerophsopholipids	Lipid	0.029(0.005)	1.36E-07	0.057(0.020)	7.68E-03	0.030(0.005)	3.36E-09	0.013
Cream	Hydroxysphingomyeline C14:1	Sphingolipids	Lipid	0.034(0.005)	8.09E-10	0.039(0.016)	2.37E-02	0.035(0.005)	2.22E-11	0.013
Cream	lysoPhosphatidylcholine acyl C28:1	Glycerophsopholipids	Lipid	0.025(0.005)	7.57E-07	0.050(0.025)	5.63E-02	0.026(0.005)	1.17E-07	0.012
Herbal tea	Phosphatidylcholine acyl-alkyl C38:3	Glycerophsopholipids	Lipid	-0.012(0.002)	1.78E-08	-0.001(0.005)	7.75E-01	-0.011(0.002)	9.91E-08	0.017
Herbal tea	Phosphatidylcholine acyl-alkyl C40:4	Glycerophsopholipids	Lipid	-0.010(0.002)	6.94E-07	-0.003(0.006)	5.44E-01	-0.009(0.002)	8.01E-07	0.014
Herbal tea	Phosphatidylcholine acyl-alkyl C40:3	Glycerophsopholipids	Lipid	-0.014(0.002)	3.64E-09	-0.004(0.006)	5.40E-01	-0.013(0.002)	5.57E-09	0.013
Herbal tea	Phosphatidylcholine acyl-alkyl C38:2	Glycerophsopholipids	Lipid	-0.018(0.003)	1.61E-07	-0.002(0.008)	8.05E-01	-0.016(0.003)	6.68E-07	0.012
Herbal tea	Phosphatidylcholine diacyl C42:4	Glycerophsopholipids	Lipid	-0.014(0.003)	1.51E-07	-0.002(0.008)	7.43E-01	-0.013(0.002)	2.80E-07	0.011
Herbal tea	Phosphatidylcholine acyl-alkyl C38:1	Glycerophsopholipids	Lipid	-0.020(0.003)	1.65E-08	-0.001(0.010)	9.33E-01	-0.018(0.003)	6.19E-08	0.010
Oily fish	Phosphatidylcholine diacyl C40:6	Glycerophsopholipids	Lipid	0.073(0.011)	1.07E-11	0.015(0.021)	4.88E-01	0.061(0.009)	7.34E-11	0.041
Oily fish	Phosphatidylcholine diacyl C38:6	Glycerophsopholipids	Lipid	0.077(0.011)	9.80E-13	0.012(0.021)	5.68E-01	0.064(0.009)	1.04E-11	0.041
White/brown bread, refined grains	Octenoylcarnitine	Acylcarnitines	Lipid	0.013(0.002)	1.32E-08	0.008(0.005)	8.23E-02	0.012(0.002)	2.45E-09	0.031
Wine	Phosphatidylcholine diacyl C36:5	Glycerophsopholipids	Lipid	0.018(0.003)	1.16E-08	0.011(0.008)	1.73E-01	0.017(0.003)	3.63E-09	0.038
Wine	Phosphatidylcholine diacyl C32:1	Glycerophsopholipids	Lipid	0.015(0.003)	2.83E-07	0.013(0.011)	2.25E-01	0.015(0.003)	9.27E-08	0.028

Table 6. List of metabolites associated with food intake from the Biocrates platform

Notes: Table shows results of the linear regression analysis for the discovery population (excluding monozygotic twins discordant for each food group intake), the MZ discordant twin sample and the fixed effects meta-analysis of both groups. Only significant associations are shown which includes those associations passing the bonferroni cut-off in the discovery and fixed effects analyses (1.08x10⁻⁶ = 0.05/[77 food variables x 601 detected metabolites]) and in the same direction in the discordant twin group. The R-squared was calculated for the relevant metabolite and food group in the study population. The Biocrates platform is a targeted platform which measured 163 metabolites in blood; data were available for 858 twins.

						-	Previous
Pathway	Subpathway	Motabolito	Poplicated	Novol	Riological rolo	Previous dietary	disease/metabolic
Lip	Essential fatty acid	Docosahexaenoat e (DHA; 22:6n3)	↑ Oily fish ↑ Other seafood	↑ Wine ↓ Baked goods	Essential fatty acid, derived endogenously from α -linolenic acid and in high concentrations in oily fish.	 ↑ Fish (excluding shellfish); ↑ Other vegetables: Celery, green beans, squash, cucumbers; ↑ White rice; ↓ Chips (Guertin et al., 2014) ↑ Fish and seafood (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) 	 ↓ NASH (Kalhan et al., 2011) ↓ Simvastatin treatment (Chen et al., 2011)
Lip	Essential fatty acid	Docosapentaenoat e (n3 DPA; 22:5n3)	↑ Wine	↑ Other seafood ↑ Oily fish	Essential fatty acid, derived endogenously from omega-3 and 6 fatty acids and in high concentrations in oily fish.	 ↑ Eggs (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) ↑ Alcohol (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) 	
Lip	Essential fatty acid	Eicosapentaenoate (EPA; 20:5n3)	↑ Oily fish ↑ Other seafood	 ↑ Wine ↓ Baked goods ↑ Avocado ↑ High fat salad dressings 	Essential fatty acid, derived endogenously from α -linolenic acid and in high concentrations in oily fish.	↑ Fish and seafood (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014)	↓ NASH (Kalhan et al., 2011)
Lip	Lysolipid	1- Docosahexaenoylg lycerophosphocholi ne*	↑ Oily fish ↑ Other seafood	↑ Avocado ↑ Green leafy vegetables	Downstream product of omega-3 essential fatty acid metabolism.	↑ Fish (excluding shellfish), ↑ Healthy eating index (Guertin et al., 2014)	
Lip	Lysolipid	1- docosahexaenoylgl ycerophosphoetha nolamine		↑ Wine	Downstream product of omega-3 essential fatty acid metabolism.		
Lip	Lysolipid	1- Arachidonoylglycer ophosphoethanola mine*		↓ Oily fish	Downstream product of omega-6 essential fatty acid metabolism. Lysolipids form cellular lipid bilayer, when cleaved by lipoprotein-associated phospholipase A2 form free lysophosphatidylcholines involved in inflammatory processes and may influence artherosclerotic plaque inflammation (Goncalves et al., 2012).	↓ HEI (Guertin et al., 2014)	
Lip	Lysolipid	1-Eicosatrienoyl glycerophosphoch oline*		↓ Oily fish ↓ Other seafood			↓ Dilated cardiomyopathy (Alexander, Lombardi, Rodriguez, Mitchell, & Marian, 2011)

Table 7. Information for metabolites associated to food intakes, including biological role, previous dietary and disease associations

							Previous
Dethurses	Ordenation		Devilianted	Nevel	Distantiant	Previous dietary	disease/metabolic
Pathway	Subpathway		Replicated	Novel	Biological role		associations
Lip	Lysolipid	1- Linoleoylglyceroph osphoethanolamin e*		↓ Oily fish		↓ HEI (Guertin et al., 2014)	↓ Dilated cardiomyopathy (Alexander et al., 2011)
Lip	Lysolipid	1- Oleoylglycerophos phoethanolamine		↓ Oily fish			↓ Dilated cardiomyopathy (Alexander et al., 2011)
Lip	Fatty acid, dicarboxylate	3-Carboxy-4- methyl-5-propyl-2- furanpropanoate (CMPF)	↑ Oily fish ↑ Other seafood	↑ Avocado ↑ Green leafy vegetables	Furan fatty acid, found in significant concentrations in fish. Thought to be a uremic toxin (Miyamoto et al., 2012). Though to induce beta cell dysfunction (Prentice et al., 2014).	 ↑ Fish and shellfish; ↑ Alliums (garlic and onions); ↑ Greens: lettuce, spinach, green pepper (Guertin et al., 2014) ↑ Fish and seafood (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) ↓ Dietary choline depletion (Sha et al., 2010) 	
Lip	Long chain fatty acid	Stearidonate (18:4n3)	↑ Wine		Derived endogenously from essential fatty acid α-linolenic acid.	↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014)	
Lip	Long chain fatty acid	10-Nonadecenoate (19:1n9)		↑ Butter	Monounsaturate of nonadecenoate (19:0).	↓ Sugar-rich foods and beverages, dietary sucrose and carbohydrate (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014)	
Lip	Long chain fatty acid	Myristate (14:0)		↑ Butter	Contained in most animal and vegetable fats. Higher concentrations in plasma associated with heart failure (Yamagishi, Nettleton, & Folsom, 2008).	↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014)	↓ T2D, IFG (Menni et al., 2013)
Lip	Long chain fatty acid	Pentadecanoate (15:0)	↑ Butter		Produced by ruminal bacteria in livestock, found in highest concentrations in dairy products, butter and meats from ruminant animals.	↑ Butter (Guertin et al., 2014)	↓ T2D, IFG (Menni et al., 2013)
Lip	Medium chain fatty acid	10-Undecenoate (11:1n1)	↑ Butter		Monounsaturated fatty acid found in oils and animal fats and produced endogenously.	↑ Butter (57)	↓ NASH, steatosis (51)
Lip	Fatty acid, branched	15-Methylpalmitate (isobar with 2- methylpalmitate)	↑ Butter		Methylated long chain fatty acid, found to be anti-inflammatory and anti-fibrotic in liver disease animal models (Mantawy, Tadros, Awad, Hassan, & El- Demerdash, 2012).	↑ Butter, ↓ HEI (Guertin et al., 2014)	↓ T2D, IFG (Menni et al., 2013)

Table 7. Information for metabolites associated to food intakes, including biological role, previous dietary and disease associations

							Previous
Pathway	Subpathway	Metabolite	Replicated	Novel	Biological role	Previous dietary	disease/metabolic
Lip	Medium chain fatty acid	Caprate (10:0)	Replicated	↑ Wine	Saturated fatty acid found in oils and animal fats and produced endogenously.		
Lip	Medium chain fatty acid	Caprylate (8:0)		↑ Wine	Saturated fatty acid found in oils and animal fats and produced endogenously.		
Lip	Phosphatidyl- choline	lysoPhosphatidylch oline acyls C17:0 & C28:1		↑ Cream			
Lip	Phosphatidyl- choline	Phosphatidylcholin e acyl-alkyls C38:1, C38:2, C38:3, C40:3 & C40:4		↓ Herbal tea			
Lip	Phosphatidyl- choline	Phosphatidylcholin e diacyls C32:1 & C36:5		↑ Wine		 ↑ Factor 1: ↑ Fish and poultry, ↓ Confectionary, cake, cookies, desserts, margarine, tea, whole grain bread, pasta, rice and high fat cheese; ↑ Factor 2: ↑ sauce and butter, ↓ fish, whole grain bread, tea, grain flakes, muesli (Floegel et al., 2013) 	
Lip	Phosphatidyl- choline	Phosphatidylcholin e diacyls C38:6 & C40:6		↑ Oily fish		↑ Factor 1: ↑ Fish and poultry, ↓ Confectionary, cake, cookies, desserts, margarine, tea, whole grain bread, pasta, rice and high fat cheese; ↓ Factor 2: ↑ sauce and butter, ↓ fish, whole grain bread, tea, grain flakes, muesli (Floegel et al., 2013)	
Lip	Phosphatidyl- choline	Phosphatidylcholin e diacyl C42:4		↓ Herbal tea			
Lip	Sphingomyelin	Hydroxysphingomy eline C14:1		↑ Cream		↑ Factor 1: ↑ Butter, garlic and coffee, ↓ Margarine, fresh fruit and soup; ↓ Factor 2: ↓ Butter, sweet bread spreads, high fat cheese, fresh fruit, whole grain bread, desserts, cake, cookies, high-fat dairy products (Floegel et al., 2013)	

Table 7. Information for metabolites associated to food intakes, including biological role, previous dietary and disease associations

							Previous
Pathway	Subpathway	Metabolite	Replicated	Novel	Biological role	Previous dietary	disease/metabolic
Lip	Carnitine metabolism	Nonanoylcarnitine*	Replicated	↑ Butter	Ester of carnitine with pelargonic acid (C9).		
Lip	Carnitine metabolism	Octenoylcarnitine		↑ White and brown bread, refined grains	Acylcarnitines consist of carnitine and fatty acid derivatives; formed from mitochondrial beta- oxidation.	 ↓ Factor 1: ↑ Butter, ↓ Margarine and low fat cheese; ↑ Factor 2: ↑ Cornflakes and crisps, ↓ Fish, other vegetable fat, whole grain bread, cooked vegetables, garlic, nuts, tea, cabbage, sweet bread spreads, cake, cookies, high fat cheese (Floegel et al., 2013); 	
Lip	Sterol, Steroid	4-Androsten- 3beta,17beta-diol disulfate 1*	↑ Wine ↑ Spirits and liquors		Sex steroid. Found to be a testosterone precursor and androgen receptor agonist (Chen et al., 2004).	 ↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014) ↓ Nuts and peanut butter, Sugar-rich foods and beverages, dietary sucrose and carbohydrate (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) ↑ Total alcohol (Guertin et al., 2014) 	
Lip	Sterol, Steroid	5-Alpha-androstan- 3beta,17beta-diol disulfate	↑ Wine		Sex steroid. Arrests growth of prostate cells in mice (Weihua, Lathe, Warner, & Gustafsson, 2002), promotes growth of breast cancer cells <i>in vitro</i> (Couture, Theriault, Simard, & Labrie, 1993).	 ↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014) ↓ Nuts and peanut butter, Sugar-rich foods and beverages, dietary sucrose and carbohydrate (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) ↑ Total alcohol (Guertin et al., 2014) 	
Lip	Sterol, Steroid	Epiandrosterone sulfate		↑ Wine	Sex steroid, also known as 3beta-hydroxy-5alpha- androstan-17-one sulfate. Downstream metabolite of dehydroepiandrosterone (DHEA).		
Lip	Inositol metabolism	Myo-inositol		↑ Wine	Contained in citrus fruits (Katz et al., 2011), beans, grains and nuts (Clements & Darnell, 1980).	↓ Dietary choline depletion (Sha et al., 2010)	 ↑ Insulin sensitivity (Gall et al., 2010) ↑ Dilated cardiomyopathy (Alexander et al., 2011)

Table 7. Information for metabolites associated to food intakes, including biological role, previous dietary and disease associations
							Previous
						Previous dietary	disease/metabolic
Pathway	Subpathway	Metabolite	Replicated	Novel	Biological role	associations	associations
Lip	Inositol metabolism	Scyllo-inositol	↑ Wine	↓ Baked goods ↓ Fried fish	Contained in citrus juices and grapes; has been used as a potential Alzheimer's treatment (Ma, Thomason, & McLaurin, 2012).	 ↑ Citrus fruit: oranges, orange juice, grapefruit; ↑ Wine (Guertin et al., 2014) ↑ Fruit juice (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) 	
AA	Butanoate metabolism	2-Aminobutyrate	↑ Wine		Part of AHB-related pathway. AHB is a by-product of hepatic glutathione synthesis under oxidative stress (Lord & Bralley, 2008).	 ↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014) ↓ Sugar-rich foods and beverages, dietary sucrose and carbohydrate (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) 	
AA	Cysteine, methionine, SAM, taurine metabolism	2-Hydroxybutyrate (AHB)	↑ Wine		Part of AHB-related pathway. AHB is a by-product of hepatic glutathione synthesis under oxidative stress (Lord & Bralley, 2008).	 ↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014) ↓ Sugar-rich foods and beverages, dietary sucrose and carbohydrate (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) 	↑ T2D, IFG (Menni et al., 2013) ↓ Insulin sensitivity (Gall et al., 2010) ↑ BMI (Moore, 2013) ↑ Dilated cardiomyopathy (Alexander et al., 2011)
AA	Valine, leucine and isoleucine metabolism	alpha- Hydroxyisovalerate	↑ Wine		Part of AHB-related pathway. AHB is a by-product of hepatic glutathione synthesis under oxidative stress (Lord & Bralley, 2008).	 ↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014) ↓ Sugar-rich foods and beverages, dietary sucrose and carbohydrate (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) 	 ↑ Dilated cardiomyopathy (Alexander et al., 2011) ↑ 28-day mortality in critically ill patients (Rogers et al., 2014)
AA	Valine, leucine and isoleucine metabolism	beta- Hydroxyisovalerate		↑ Wine	Part of AHB-related pathway. AHB is a by-product of hepatic glutathione synthesis under oxidative stress (Lord & Bralley, 2008).		
AA	Valine, leucine and isoleucine metabolism	3-Methyl-2- oxobutyrate		↑ Wine	Derived from branched chain amino acid, precursor to leucine and valine synthesis.		↑ T2D, IFG (Menni et al., 2013) ↑ BMI (Moore, 2013)
AA	Valine, leucine and isoleucine metabolism	4-Methyl-2- oxopentanoate		↑ Wine	Derived from branched chain amino acid, leucine, metabolism; accumulates, and may contribute to DNA damage and neurotoxicity, in maple-syrup urine disease (Mescka et al., 2014).		↑ T2D, IFG (Menni et al., 2013) ↑ BMI (Moore, 2013)

Table 7. Information for	metabolites associated t	o food intakes, in	ncluding biologica	al role, previous dietary	/ and disease associations

Pathway	Subpathway	Metabolite	Replicated	Novel	Biological role	Previous dietary associations	Previous disease/metabolic associations
AA	Lysine metabolism	Pipecolate		↑ Wine ↓ Confectionary & jams	Present in citrus fruits (Servillo et al., 2012). Thought to be formed primarily by gut bacterial lysine degradation (Fujita, Fujita, Kodama, Hada, & Higashino, 2003).		
AA	Phenylalanine & tyrosine metabolism	3-(4- Hydroxyphenyl)lact ate		↑ Wine	The L-form is produced from tyrosine metabolism. The unusual D-form is of bacterial origin (Spaapen, Ketting, Wadman, Bruinvis, & Duran, 1987), particularly by Bifidobacteria and lactobacilli, from dietary phenolic compounds and has shown to reduce mitochondrial and neutrophil ROS production (Beloborodova et al., 2012).		↑ BMI (Moore, 2013)
AA	Phenylalanine & tyrosine metabolism	3- Phenylpropionate (hydrocinnamate)		↓ Fried fish ↓ Savoury pies ↑ Apples & pears	Formed by gut bacterial degradation of tyrosine (Smith & Macfarlane, 1996) and phenolic compounds (Anson et al., 2009; van Dorsten et al., 2012).		↑ NASH vitamin E responders at baseline (62)
AA	Phenylalanine & tyrosine metabolism	3,4- dihydroxyphenylac etate sulfate		↓ Other seafood			
AA	Tryptophan metabolism	Indolepropionate		↑ Bananas ↑ Apples & pears	Formed by gut bacterial degradation of tryptophan (Smith & Macfarlane, 1996). Acts as an antioxidant (Karbownik et al., 2001).	↓ Eggs, ↓ Red meat (Guertin et al., 2014) ↓ Dietary choline depletion (Sha et al., 2010)	 ↑ Insulin sensitivity (Gall et al., 2010) ↑ NASH vitamin E responders at baseline (Cheng, Joyce, Yates, Aouizerat, & Sanyal, 2012) ↓ Dilated cardiomyopathy (Alexander et al., 2011) ↓ Muscle mass indices in elderly (Lustgarten, Price, Chale, Phillips, & Fielding, 2014)
AA	Tryptophan metabolism	Tryptophan betaine		↑ Allium vegetables	Indole alkaloid found in legumes of <i>Erythrina</i> species, also called hypaphorine. Found to induce sleep in mice (Ozawa, Honda, Nakai, Kishida, & Ohsaki, 2008).	 ↑ Peanuts (Guertin et al., 2014) ↑ Nuts and peanut butter (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) 	

	Table 7. Information for metabolites	associated to food intakes	. including biological r	role, previous dietar	v and disease associations
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							Previous
Dethurou	Submothway	Motobolito	Deplicated	Nevel	Dielegical role	Previous dietary	disease/metabolic
AA	Creatine metabolism	Creatine	Replicated	↑ Meat ↑ Poultry	Red meat a major source; vegetarians have lower circulating levels (Delanghe et al., 1989).	Sociations Sugar-rich foods and beverages, dietary sucrose and carbohydrate (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014)	Insulin sensitivity (Gall et al., 2010) Steatosis vs. NASH (Kalhan et al., 2011) Dilated cardiomyopathy (Alexander et al., 2011)
AA	Glutamate metabolism	Pyroglutamine*	↓ Poultry	↓ Other seafood ↓ Meat		↓ Poultry (chicken) (Guertin et al., 2014) ↑ Dietary choline depletion (Sha et al., 2010)	 ↑ Active TB (Weiner et al., 2012) ↑ Dilated cardiomyopathy (Alexander et al., 2011)
AA	Urea cycle; arginine-, proline-, metabolism	Trans-4- hydroxyproline		↑ Meat ↑ Processed meat ↑ Beef burgers	Major component of collagen; circulating levels increase following oral gelatin ingestion (Ohara, Matsumoto, Ito, Iwai, & Sato, 2007).	↑ Dietary choline depletion (Sha et al., 2010)	
Xeno	Xanthine metabolism	1-Methylxanthine	↑ Coffee			↑ Coffee (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) (Guertin et al., 2015) ↑ β-carotene supplementation in smokers (Mondul et al., 2013)	
Xeno	Xanthine metabolism	7-Methylxanthine		↑ Chocolate		↑ Coffee (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) ↑ Chocolate (Guertin et al., 2014)	
Xeno	Xanthine metabolism	Theobromine	↑ Chocolate		Bitter alkaloid characteristic of the cacao plant.	 ↑ Sugar-rich foods and beverages, dietary sucrose and carbohydrate (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) ↑ Coffee (Guertin et al., 2015) 	
Xeno	Xanthine metabolism	Theophylline		↑ Wine	Theophylline is methylxanthine metabolite found in caffeinated beverages and foods but also administered as a treatment for COPD and asthma. Alcohol has been shown to delay plasma clearance of theophylline (Thompson, 1992).		
Xeno	Benzoate	3-methyl catechol	↑ Coffee	↓ Black tea		↑ Coffee (Guertin et al.,	

Table 7. Information for metabolites associated to food intakes, including biological role, previous dietary and disease associations

						Previous dietarv	Previous disease/metabolic
Pathway	Subpathway	Metabolite	Replicated	Novel	Biological role	associations	associations
	metabolism	sulfate 1				2015)	
Xeno	Benzoate metabolism	Catechol sulfate	↑ Coffee		Chlorogenic acid derived metabolite (Lang et al., 2013).	 ↑ Coffee (Guertin et al., 2014) ↑ β-carotene supplementation in smokers (Mondul et al., 2013) 	↑ Insulin sensitivity (Gall et al., 2010)
Xeno	Benzoate metabolism	O-methyl catechol sulfate		↑ Coffee			
Xeno	Benzoate metabolism	4- Ethylphenylsulfate		↑ Soy and other milks	Synthesized by gut bacteria, elevated in chronic kidney disease (Itoh, Ezawa, Kikuchi, Tsuruta, & Niwa, 2013); elevated levels may indicate intestinal permeability (Hsiao et al., 2013).	↑ Tofu (Guertin et al., 2014)	
Xeno	Benzoate metabolism	Benzoate		↑ Wine	Added to foods to prevent mold and bacterial growth (often as sodium benzoate in wine).		↓ BMI (Moore, 2013)
Xeno	Chemical	3-hydroxypyridine sulfate	↑ Coffee	↓ Black tea		↑ Coffee (Guertin et al., 2015)	
Xeno	Chemical	2-aminophenol sulfate		↑ Porridge	Benzoxazinoid metabolite excreted in urine following whole grain rye bread consumption (Bondia-Pons et al., 2013).		
Xeno	Food component, Plant	Quinate	↑ Coffee	↓ Black tea	Phenolic product of chlorogenic acid, acts as an antioxidant (Lang et al., 2013).	 ↑ Coffee (Guertin et al., 2014; Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) ↓ Sugar-sweetened beverages: soda, fruit punch (Guertin et al., 2014) 	↑ Insulin sensitivity
Xeno	Food component, Plant	Piperine	↑ Wine		Compound in black pepper with antioxidant effects (Srinivasan, 2014).	↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014)	
Xeno	Food component, Plant	Ergothioneine		↑ Mushroom ↓ Savoury pies	Thiol compound which enters human cells via the organic cation transporter OCTN1. Is found in particularly high concentrations in specialty mushrooms, oat bran and beans and demonstrates protection against copper(II)-induced toxicity (Ey, Schomig, & Taubert, 2007).		↓ Parkinson's disease patients (Ey et al., 2007)

Table 7. Information for metabolites associated to food intakes, including biological role, previous dietary and disease associations

Pathway	Subpathway	Metabolite	Replicated	Novel	Biological role	Previous dietary associations	Previous disease/metabolic associations
Xeno	Food component, Plant	Stachydrine	↑ Fruit juice ↑ Citrus fruit		Present in citrus fruits and juices (Servillo, Giovane, Balestrieri, Cautela, & Castaldo, 2011).	 ↑ Citrus fruit: oranges, orange juice, grapefruit (Guertin et al., 2014) ↑ Fruit juice (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) ↓ Dietary choline depletion (Sha et al., 2010) 	↑ Dilated cardiomyopathy (Alexander et al., 2011)
Cho	Glycolysis, gluconeogenes is, pyruvate metabolism	Glycerate	↑ Citrus fruit	↓ Confectionary & jams ↑ Tomatoes	Contained in citrus fruits (Katz et al., 2011)	 ↑ Fruits and vegetables; ↑ Fruit juice (Zheng, Yu, Alexander, Steffen, & Boerwinkle, 2014) 	↑ Insulin sensitivity (Gall et al., 2010)
Cho	Glycolysis, gluconeogenes is, pyruvate metabolism	1,5-Anhydroglucitol (1,5-AG)		↑ Other seafood	Marker of glycaemic control and of dietary origin (Yamanouchi et al., 1989).	↓ β-carotene supplementation in smokers (Mondul et al., 2013)	↓ T2D, IFG (Menni et al., 2013) ↓ BMI (Moore, 2013)
Cho	Nucleotide sugars, pentose metabolism	Threitol		↑ Apples and pears	Primary end product of D-xylose metabolism. Found in the edible fungus Armillaria mellea, jute and the pigeon pea plant FDB002261.		
Vit	Vitamin B6 metabolism	Pyridoxate		↑ High fibre breakfast cereals	Essential nutrient, coenzyme for synthesis of amino acids, neurotransmitters (serotonin, norepinephrine), sphingolipids, and aminolevulinic acid.	 ↑ Vitamins/supplements; ↑ Healthy eating index; ↑ Other fruits: plums, apricots, peaches, prunes, raisins, grapes, pineapple (Guertin et al., 2014) ↓ Dietary choline depletion (Sha et al., 2010) 	
Рер	Dipeptide	Cyclo(leu-pro)		↑ Coffee ↓ Black tea	Cyclo(leu-pro) is a diketopiperazine which contributes to the bitter flavor of roasted coffee beans (Ginz & Engelhardt, 2000).	↓ Starchy vegetables: white potatoes, corn, and peas, ↑ Total alcohol (Guertin et al., 2014) ↑ Alcohol (Zheng, Yu, Alexander, Steffen, Nettleton, et al., 2014)	

Table 7 Information for metabolites associated to food intakes	s including biological rol	e previous dietary a	ad disease associations
Table 7. Information for metabolites associated to food intakes	s, including biological for	e, previous uletary al	iu uisease associations

Notes: Associations are reported which replicate previous findings from food intake and blood metabolite association studies and those which associations which we believe to be novel. Biological roles of most metabolites are defined and some previous disease associations with the appropriate metabolite in blood are included.

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				Discovery		Discor	dant	Meta-analysis		Unique
Variable	Metabolite name	Pathway	Super- pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	to pattern
Mediterranean Diet Score	X-11315			0.117(0.010)	3.22E-28	0.085(0.018)	2.01E-06	0.109(0.009)	7.94E-34	No
Mediterranean Diet Score	X-11469			0.104(0.010)	3.86E-23	0.091(0.016)	3.08E-08	0.100(0.009)	9.16E-31	No
Mediterranean Diet Score	X-02269			0.101(0.010)	7.10E-22	0.093(0.016)	2.98E-08	0.099(0.009)	1.70E-29	No
Mediterranean Diet Score	docosahexaenoate (DHA: 22:6n3)	Essential fatty acid	Lipid	0.100(0.010)	5.78E-21	0.090(0.017)	3.88E-07	0.097(0.009)	2.45E-27	No
Mediterranean Diet Score	3-carboxy-4-methyl-5- propyl-2- furanpropanoate (CMPF)	Fatty acid, dicarboxylate	Lipid	0.084(0.010)	4.57E-17	0.082(0.016)	2.72E-07	0.084(0.008)	1.82E-23	No
Mediterranean Diet Score	eicosapentaenoate (EPA: 20:5n3)	Essential fatty acid	Lipid	0.089(0.010)	2.82E-18	0.062(0.019)	9.62E-04	0.083(0.009)	9.54E-21	No
Mediterranean Diet Score	1- docosahexaenoylglyce rophosphocholine*	Lysolipid	Lipid	0.084(0.011)	2.16E-15	0.068(0.018)	2.11E-04	0.080(0.009)	1.28E-18	No
Mediterranean Diet Score	indolepropionate	Tryptophan metabolism	Amino acid	0.086(0.011)	7.59E-16	0.054(0.018)	3.44E-03	0.078(0.009)	1.60E-17	No
Mediterranean Diet Score	X-21365 [trimethyl-N- aminovalerate]			-0.076(0.011)	1.60E-12	-0.064(0.018)	3.20E-04	-0.073(0.009)	1.48E-15	No
Mediterranean Diet Score	tryptophan betaine	Tryptophan metabolism	Amino acid	0.089(0.012)	2.36E-13	0.068(0.022)	2.53E-03	0.084(0.011)	1.65E-15	No
Mediterranean Diet Score	X-11847			0.091(0.013)	9.97E-13	0.074(0.024)	2.38E-03	0.088(0.011)	6.32E-15	Yes
Mediterranean Diet Score	X-11381			-0.075(0.011)	5.75E-12	-0.052(0.018)	3.27E-03	-0.069(0.009)	8.27E-14	No
Mediterranean Diet Score	X-11849			0.094(0.014)	3.05E-11	0.067(0.023)	3.69E-03	0.086(0.012)	4.15E-13	Yes
Mediterranean Diet Score	X-12798			-0.067(0.011)	2.68E-09	-0.076(0.019)	8.03E-05	-0.069(0.010)	6.80E-13	No
Mediterranean Diet Score	ergothioneine	Food component, Plant	Xenobiotics	0.083(0.016)	1.08E-07	0.107(0.026)	4.21E-05	0.090(0.013)	1.56E-11	No
Mediterranean Diet Score	X-13477			0.060(0.012)	4.66E-07	0.086(0.021)	6.66E-05	0.067(0.010)	1.56E-10	Yes
Mediterranean Diet Score	scyllo-inositol	Inositol metabolism	Lipid	0.067(0.012)	1.52E-08	0.061(0.022)	5.19E-03	0.066(0.010)	2.25E-10	No
Mediterranean Diet Score	piperine	Food component, Plant	Xenobiotics	0.060(0.012)	4.12E-07	0.050(0.020)	1.39E-02	0.057(0.010)	1.73E-08	No
Fruit & Vegetable	X-11315	1 Idin		0.145(0.009)	5.48E-52	0.130(0.022)	1.20E-08	0.142(0.008)	3.87E-63	No
Fruit & Vegetable	X-11469			0.112(0.010)	3.63E-29	0.083(0.018)	1.16E-05	0.106(0.009)	3.39E-34	No
Fruit & Vegetable	X-11372			-0.109(0.009)	6.97E-32	-0.061(0.021)	4.25E-03	-0.102(0.008)	5.27E-34	No

			Supor	Discovery		Discordant		Meta-analysis		Unique
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	to pattern
Fruit & Vegetable	X-02269		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	0.108(0.010)	1.21E-26	0.079(0.019)	5.22E-05	0.102(0.009)	7.46E-31	No
Fruit & Vegetable	glycerate	Glycolysis, gluconeogenesis, pyruvate metabolism	Carbo- hydrate	0.094(0.010)	1.57E-21	0.094(0.019)	1.78E-06	0.094(0.009)	1.97E-27	No
Fruit & Vegetable	eicosapentaenoate (EPA; 20:5n3)	Essential fatty acid	Lipid	0.105(0.011)	2.63E-22	0.069(0.018)	1.66E-04	0.095(0.009)	1.86E-25	No
Fruit & Vegetable	scyllo-inositol	Inositol metabolism	Lipid	0.093(0.010)	1.08E-20	0.068(0.021)	1.08E-03	0.088(0.009)	2.22E-23	No
Fruit & Vegetable	ergothioneine	Food component, Plant	Xenobiotics	0.117(0.013)	7.72E-20	0.080(0.030)	8.94E-03	0.111(0.012)	8.16E-22	No
Fruit & Vegetable	docosahexaenoate (DHA; 22:6n3)	Essential fatty acid	Lipid	0.102(0.012)	5.49E-18	0.067(0.018)	1.91E-04	0.091(0.010)	6.05E-21	No
Fruit & Vegetable	threonate	Ascorbate and aldarate metabolism	Cofactors and vitamins	0.080(0.009)	5.50E-17	0.090(0.022)	6.95E-05	0.081(0.009)	6.46E-21	Yes
Fruit & Vegetable	indolepropionate	Tryptophan metabolism	Amino acid	0.081(0.010)	1.81E-16	0.085(0.020)	3.00E-05	0.082(0.009)	8.42E-21	No
Fruit & Vegetable	1- docosahexaenoylglyce rophosphocholine*	Lysolipid	Lipid	0.077(0.009)	8.09E-18	0.048(0.020)	2.03E-02	0.072(0.008)	5.51E-19	No
Fruit & Vegetable	3-carboxy-4-methyl-5- propyl-2- furanpropanoate (CMPF)	Fatty acid, dicarboxylate	Lipid	0.082(0.010)	9.19E-17	0.053(0.020)	7.55E-03	0.076(0.009)	2.74E-18	No
Fruit & Vegetable	tryptophan betaine	Tryptophan metabolism	Amino acid	0.083(0.010)	9.57E-16	0.064(0.025)	1.06E-02	0.080(0.009)	1.94E-17	No
Fruit & Vegetable	X-14056			-0.071(0.010)	2.40E-13	-0.081(0.021)	1.29E-04	-0.073(0.009)	6.78E-17	Yes
Fruit & Vegetable	X-12056			0.094(0.012)	1.19E-14	0.086(0.029)	3.95E-03	0.093(0.011)	6.91E-17	Yes
Fruit & Vegetable	1,5-anhydroglucitol (1,5-AG)	Glycolysis, gluconeogenesis, pyruvate metabolism	Carbo- hydrate	-0.074(0.009)	2.92E-15	-0.054(0.021)	1.32E-02	-0.071(0.009)	1.02E-16	No
Fruit & Vegetable	3-phenylpropionate (hydrocinnamate)	Phenylalanine & tyrosine metabolism	Amino acid	0.077(0.011)	4.84E-12	0.089(0.020)	1.72E-05	0.080(0.010)	1.73E-16	No
Fruit & Vegetable	X-11847			0.077(0.011)	2.85E-11	0.092(0.025)	3.65E-04	0.080(0.010)	2.68E-14	Yes
Fruit & Vegetable	X-11849			0.085(0.013)	1.96E-11	0.083(0.027)	2.38E-03	0.085(0.011)	9.86E-14	Yes
Fruit & Vegetable	pipecolate	Lysine metabolism	Amino acid	0.060(0.009)	6.41E-11	0.057(0.020)	3.94E-03	0.059(0.008)	6.30E-13	No
Fruit & Vegetable	hippurate	Benzoate metabolism	Xenobiotics	0.061(0.009)	2.42E-11	0.042(0.021)	4.86E-02	0.058(0.008)	3.29E-12	Yes

			Super	Discovery		Discordant		Meta-analysis		Unique
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	to pattern
Fruit & Vegetable	X-09789	, , , , , , , , , , , , , , , , , , ,		0.062(0.010)	7.79E-10	0.059(0.025)	1.84E-02	0.061(0.009)	3.50E-11	No
Fruit & Vegetable	catechol sulfate	Benzoate metabolism	Xenobiotics	0.058(0.010)	1.78E-08	0.071(0.021)	7.28E-04	0.060(0.009)	4.21E-11	No
Fruit & Vegetable	myo-inositol	Inositol metabolism	Lipid	0.051(0.009)	5.14E-09	0.056(0.019)	3.08E-03	0.052(0.008)	4.43E-11	No
Fruit & Vegetable	threitol	Nucleotide sugars, pentose metabolism	Carbo- hydrate	0.056(0.010)	6.64E-09	0.058(0.021)	6.08E-03	0.056(0.009)	1.07E-10	No
Fruit & Vegetable	X-11261			-0.062(0.011)	2.39E-08	-0.057(0.019)	3.08E-03	-0.061(0.010)	2.08E-10	Yes
Fruit & Vegetable	X-12063			-0.047(0.009)	1.73E-07	-0.052(0.018)	4.69E-03	-0.048(0.008)	2.31E-09	Yes
Fruit & Vegetable	X-11858			0.080(0.015)	9.59E-08	0.077(0.034)	2.48E-02	0.080(0.014)	5.28E-09	No
Fruit & Vegetable	proline	Urea cycle; arginine-, proline-, metabolism	Amino acid	-0.053(0.010)	2.60E-07	-0.060(0.027)	2.87E-02	-0.054(0.010)	1.95E-08	Yes
Fruit & Vegetable	N-acetylornithine	Urea cycle; arginine-, proline-, metabolism	Amino acid	0.049(0.010)	3.45E-07	0.052(0.024)	2.92E-02	0.050(0.009)	2.57E-08	Yes
Fruit & Vegetable	gamma-glutamylvaline	gamma-glutamyl	Peptide	-0.047(0.009)	1.04E-06	-0.043(0.018)	1.71E-02	-0.046(0.008)	4.79E-08	Yes
High Alcohol	alpha- hydroxyisovalerate	Valine, leucine and isoleucine metabolism	Amino acid	0.165(0.014)	1.32E-32	0.108(0.036)	3.05E-03	0.158(0.013)	2.20E-35	No
High Alcohol	X-11795			0.150(0.013)	1.93E-28	0.118(0.026)	1.02E-05	0.143(0.012)	1.19E-33	No
High Alcohol	scyllo-inositol	Inositol metabolism	Lipid	0.146(0.015)	2.89E-22	0.128(0.033)	1.54E-04	0.143(0.014)	4.20E-26	No
High Alcohol	piperine	Food component, Plant	Xenobiotics	0.133(0.015)	1.91E-18	0.084(0.026)	1.63E-03	0.121(0.013)	1.62E-20	No
High Alcohol	4-androsten- 3beta,17beta-diol disulfate 1*	Sterol, Steroid	Lipid	0.113(0.015)	9.34E-14	0.157(0.031)	1.35E-06	0.122(0.014)	3.27E-19	No
High Alcohol	X-09789			-0.118(0.014)	6.15E-16	-0.107(0.030)	4.23E-04	-0.116(0.013)	4.99E-19	No
High Alcohol	ergothioneine	Food component, Plant	Xenobiotics	0.147(0.019)	1.65E-14	0.147(0.037)	1.21E-04	0.147(0.017)	2.44E-18	No
High Alcohol	X-11799			0.116(0.015)	9.27E-14	0.137(0.032)	4.04E-05	0.120(0.014)	6.07E-18	No
High Alcohol	X-01911			0.129(0.016)	7.26E-16	0.074(0.027)	7.15E-03	0.115(0.014)	3.77E-17	No
High Alcohol	tryptophan betaine	Tryptophan metabolism	Amino acid	0.129(0.016)	6.70E-16	0.069(0.030)	2.27E-02	0.116(0.014)	1.08E-16	No
High Alcohol	X-12798			-0.103(0.014)	8.77E-13	-0.086(0.025)	6.77E-04	-0.099(0.012)	1.58E-15	No

			Super	Discovery		Discor	dant	Meta-analysis		Unique
Variable	Metabolite name	Pathway	pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	to pattern
High Alcohol	5alpha-androstan- 3beta,17beta-diol disulfate	Sterol, Steroid	Lipid	0.109(0.015)	1.02E-12	0.130(0.038)	7.44E-04	0.112(0.014)	1.73E-15	No
High Alcohol	X-21365 [trimethyl-N- aminovalerate]			-0.083(0.013)	1.72E-10	-0.123(0.026)	4.64E-06	-0.091(0.012)	3.91E-15	No
High Alcohol	gamma-tocopherol	Tocopherol metabolism	Cofactors and vitamins	0.107(0.015)	4.11E-13	0.062(0.029)	3.39E-02	0.098(0.013)	6.75E-14	Yes
High Alcohol	2-aminobutyrate	Butanoate metabolism	Amino acid	0.088(0.014)	1.23E-10	0.109(0.030)	2.85E-04	0.092(0.012)	1.08E-13	No
High Alcohol	X-10395			0.097(0.014)	3.99E-12	0.066(0.030)	2.90E-02	0.091(0.013)	3.80E-13	No
High Alcohol	X-04495			0.090(0.014)	4.83E-11	0.079(0.026)	3.16E-03	0.088(0.012)	4.06E-13	No
High Alcohol	X-11381			-0.073(0.013)	2.04E-08	-0.096(0.028)	6.04E-04	-0.077(0.012)	4.54E-11	No
High Alcohol	X-11847			0.087(0.015)	1.42E-08	0.075(0.035)	3.16E-02	0.085(0.014)	1.11E-09	Yes
High Alcohol	4-vinylphenol sulfate	Benzoate metabolism	Xenobiotics	0.078(0.014)	7.28E-08	0.074(0.027)	7.55E-03	0.077(0.013)	1.50E-09	Yes
High Alcohol	10-undecenoate (11:1n1)	Medium chain fatty acid	Lipid	0.072(0.014)	4.15E-07	0.075(0.028)	7.09E-03	0.073(0.013)	8.17E-09	No
High Alcohol	X-12816			0.093(0.017)	1.19E-07	0.075(0.036)	3.97E-02	0.089(0.016)	1.22E-08	No
High Alcohol	X-12038			0.065(0.013)	9.44E-07	0.057(0.028)	4.23E-02	0.064(0.012)	1.02E-07	No
Traditional English	stachydrine	Food component, Plant	Xenobiotics	-0.100(0.013)	1.17E-13	-0.076(0.025)	2.74E-03	-0.094(0.012)	9.94E-16	No
Traditional English	X-11315			-0.099(0.017)	4.61E-09	-0.082(0.025)	1.08E-03	-0.093(0.014)	1.68E-11	No
Traditional English	3-phenylpropionate (hydrocinnamate)	Phenylalanine & tyrosine metabolism	Amino acid	-0.083(0.016)	4.19E-07	-0.089(0.025)	4.17E-04	-0.084(0.014)	5.18E-10	No
Traditional English	X-11372			0.083(0.016)	2.83E-07	0.083(0.024)	7.93E-04	0.083(0.013)	6.68E-10	No
Traditional	creatine	Creatine	Amino acid	0.089(0.015)	6.26E-09	0.050(0.022)	2.63E-02	0.076(0.013)	1.19E-09	No
Traditional	X-11381			0.072(0.014)	1.84E-07	0.070(0.026)	7.14E-03	0.072(0.012)	3.71E-09	No
Dieting	X-14473			-0.143(0.017)	3.69E-17	-0.161(0.028)	2.78E-08	-0.148(0.014)	8.46E-25	No
Dieting	X-02249			0.097(0.016)	8.79E-10	0.072(0.029)	1.35E-02	0.091(0.014)	4.01E-11	No
Dieting	quinate	Food component, Plant	Xenobiotics	-0.098(0.017)	1.70E-08	-0.088(0.029)	2.86E-03	-0.095(0.015)	1.40E-10	No
Dieting	4-ethylphenylsulfate	Benzoate	Xenobiotics	0.104(0.019)	1.01E-07	0.093(0.039)	1.84E-02	0.102(0.017)	4.89E-09	No
Dieting	X-14374	motoboliom		-0.088(0.017)	1.27E-07	-0.065(0.031)	3.85E-02	-0.083(0.015)	1.48E-08	No
Low Meat	3-carboxy-4-methyl-5-	Fatty acid,	Lipid	-0.145(0.017)	4.24E-17	-0.139(0.034)	5.26E-05	-0.144(0.015)	3.63E-21	No

			Currer	Discov	very	Discor	dant	Meta-ana	alysis	Unique
Variable	Metabolite name propyl-2- furanpropanoate (CMPE)	Pathway dicarboxylate	super- pathway	Beta(SE)	Р	Beta(SE)	Р	Beta(SE)	Р	to pattern
Low Meat	pyroglutamine*	Glutamate	Amino acid	0.102(0.016)	1.62E-10	0.162(0.024)	1.28E-10	0.120(0.013)	8.99E-20	No
Low Meat	docosahexaenoate (DHA: 22:6n3)	Essential fatty acid	Lipid	-0.134(0.017)	4.71E-15	-0.128(0.030)	2.97E-05	-0.133(0.015)	2.65E-19	No
Low Meat	(EPA: 20:5n3)	Essential fatty acid	Lipid	-0.131(0.016)	2.89E-15	-0.102(0.026)	9.59E-05	-0.123(0.014)	8.44E-19	No
Low Meat	2-aminobutyrate	Butanoate metabolism	Amino acid	-0.114(0.017)	1.57E-11	-0.108(0.028)	1.89E-04	-0.113(0.015)	7.47E-15	No
Low Meat	X-11261	metaboliom		0.111(0.016)	1.30E-11	0.089(0.027)	9.83E-04	0.105(0.014)	4.20E-14	Yes
Low Meat	X-11469			-0.111(0.017)	2.74E-11	-0.106(0.035)	2.73E-03	-0.110(0.015)	1.88E-13	No
Low Meat	X-02269			-0.112(0.017)	2.97E-11	-0.097(0.032)	3.24E-03	-0.109(0.015)	2.61E-13	No
Low Meat	creatine	Creatine metabolism	Amino acid	-0.092(0.015)	2.80E-09	-0.117(0.028)	4.24E-05	-0.098(0.014)	4.12E-13	No
Low Meat	X-02249	metaboliom		-0.122(0.018)	6.73E-12	-0.073(0.031)	1.99E-02	-0.110(0.015)	7.70E-13	No
Low Meat	X-11372			0.112(0.017)	1.24E-10	0.076(0.024)	1.73E-03	0.099(0.014)	1.17E-12	No
Low Meat	alpha- hydroxyisovalerate	Valine, leucine and isoleucine metabolism	Amino acid	-0.106(0.016)	1.24E-10	-0.078(0.027)	4.42E-03	-0.099(0.014)	2.09E-12	No
Low Meat	15-methylpalmitate (isobar with 2- methylpalmitate)	Fatty acid, branched	Lipid	-0.109(0.019)	5.20E-09	-0.091(0.029)	2.06E-03	-0.104(0.016)	3.23E-11	No
Low Meat	X-11478			0.103(0.017)	3.48E-09	0.073(0.027)	6.92E-03	0.094(0.015)	9.86E-11	Yes
Low Meat	2-hydroxybutyrate (AHB)	Cysteine, methionine, SAM, taurine metabolism	Amino acid	-0.081(0.016)	1.70E-07	-0.095(0.028)	7.53E-04	-0.085(0.014)	4.08E-10	No
Low Meat	betaine	Glycine, serine and threonine metabolism	Amino acid	0.087(0.017)	1.90E-07	0.078(0.029)	7.05E-03	0.085(0.014)	3.87E-09	Yes
Low Meat	C-glycosyltryptophan*	Tryptophan metabolism	Amino acid	0.072(0.014)	3.27E-07	0.056(0.024)	1.77E-02	0.068(0.012)	1.80E-08	Yes
Low Meat	X-11521			0.086(0.017)	2.79E-07	0.064(0.028)	2.32E-02	0.080(0.014)	2.12E-08	Yes
Low Meat	X-11204			-0.081(0.016)	6.79E-07	-0.063(0.025)	1.12E-02	-0.076(0.014)	2.52E-08	Yes

Notes: Table shows results of the linear regression analysis for the discovery population (excluding monozygotic twins discordant for each diet pattern), the MZ discordant twin sample and the fixed effects meta-analysis of both groups. Only significant associations are shown which includes those associations passing the bonferroni cut-off in the discovery and fixed effects analyses $(1.08 \times 10^{-6} = 0.05/[77 \text{ diet phenotypes x 601 detected metabolites}])$ and passing the 5% level of significance in the discordant twin group. The Metabolon platform is a non-targeted platform which identified 456 metabolites in blood for which data were available for 3559 twins.

			Super-	Discov	ery	Discorda	int	Meta-ana	alysis	Unique to
Variable	Metabolite	Pathway	pathway	Beta(SE)	Ρ	Beta(SE)	Ρ	Beta(SE)	Ρ	patterns
Fruit & Vegetable	Octenoylcarnitine	Acylcarnitines	Lipid	-0.052(0.009)	1.96E-08	-0.021(0.018)	0.261	-0.045(0.008)	2.09E-08	No
Low Meat	Phosphatidylcholine diacyl C36:5	Glycerophospholipids	Lipid	-0.115(0.016)	8.26E-12	-0.028(0.032)	0.380	-0.096(0.015)	3.96E-11	No
Low Meat	Phosphatidylcholine diacyl C36:6	Glycerophospholipids	Lipid	-0.094(0.016)	3.81E-09	-0.024(0.036)	0.510	-0.083(0.014)	8.53E-09	Yes
Low Meat	Phosphatidylcholine acyl-alkyl C38:6	Glycerophospholipids	Lipid	-0.061(0.011)	9.29E-08	-0.016(0.023)	0.481	-0.053(0.010)	2.33E-07	Yes
Low Meat	Phosphatidylcholine diacyl C38:6	Glycerophospholipids	Lipid	-0.067(0.013)	6.80E-07	-0.037(0.035)	0.294	-0.063(0.012)	3.91E-07	No

Notes: Table shows results of the linear regression analysis for the discovery population (excluding monozygotic twins discordant for each diet pattern), the MZ discordant twin sample and the fixed effects meta-analysis of both groups. Only significant associations are shown which includes those associations passing the bonferroni cut-off in the discovery and fixed effects analyses $(1.08 \times 10^{-6} = 0.05/[77 \text{ diet phenotypes x 601 detected metabolites}])$ and in the same direction in the discordant twin group. The Biocrates platform is a targeted platform which measures 163 metabolites in blood for which data were available for 858 twins.

			Diet variable-M (Meta-analysi	letabolite s result)					SNP-Metal (Shin et al.,	bolite 2014)
Metabolite	Diet variable	Diet pattern or food	Beta(SF)	P	Gene symbol	SNP	FA/OA	FAF	Beta(SF)	P
2-Aminobutyrate	Low Meat	Pattern	-0.113(0.015)	7.47E-15	PPM1K	rs10022462	T/C	0.45	-0.012(0.002)	4.55E-11
2-Aminobutyrate	Wine	Food	0.025(0.003)	2.73E-14	SLC1A4	rs10211524	A/G	0.42	0.019(0.002)	5.59E-16
2-Aminobutyrate	Wine	Food	0.025(0.003)	2.73E-14	PPM1K	rs10022462	T/C	0.45	-0.012(0.002)	4.55E-11
2-Aminobutyrate	High Alcohol	Pattern	0.092(0.012)	1.08E-13	SLC1A4	rs10211524	A/G	0.42	0.019(0.002)	5.59E-16
3-(4- Hydroxyphenyl)lactate	Wine	Food	0.018(0.003)	1.08E-09	CCDC57	rs4625783	T/C	0.43	-0.019(0.003)	3.90E-13
3-Phenylpropionate (hydrocinnamate)	Fruit & Vegetable	Pattern	0.080(0.010)	1.73E-16	ACSM5	rs11647589	A/G	0.72	0.030(0.005)	2.85E-11
3-Phenylpropionate (hydrocinnamate)	Fried fish	Food	-0.172(0.026)	4.12E-11	ACSM5	rs11647589	A/G	0.72	0.030(0.005)	2.85E-11
3-Phenylpropionate (hydrocinnamate)	Savoury pies	Food	-0.231(0.037)	3.72E-10	ACSM5	rs11647589	A/G	0.72	0.030(0.005)	2.85E-11
3-Phenylpropionate (hydrocinnamate)	Traditional English	Pattern	-0.084(0.014)	5.18E-10	ACSM5	rs11647589	A/G	0.72	0.030(0.005)	2.85E-11
3-Phenylpropionate (hydrocinnamate)	Apples & pears	Food	0.024(0.004)	1.24E-08	ACSM5	rs11647589	A/G	0.72	0.030(0.005)	2.85E-11
alpha- Hvdroxvisovalerate	Wine	Food	0.045(0.003)	1.23E-45	HAO2	rs12141041	T/C	0.47	-0.025(0.004)	1.77E-12
alpha- Hvdroxvisovalerate	High Alcohol	Pattern	0.158(0.013)	2.20E-35	HAO2	rs12141041	T/C	0.47	-0.025(0.004)	1.77E-12
alpha- Hydroxyisovalerate	Low Meat	Pattern	-0.099(0.014)	2.09E-12	HAO2	rs12141041	T/C	0.47	-0.025(0.004)	1.77E-12
Betaine	Low Meat	Pattern	0.085(0.014)	3.87E-09	CBS	rs2851391	T/C	0.46	-0.012(0.002)	1.15E-11
Creatine	Low Meat	Pattern	-0.098(0.014)	4.12E-13	CPS1	rs715	T/C	0.71	-0.045(0.004)	9.63E-25
Creatine	Meat	Food	0.063(0.009)	8.24E-12	CPS1	rs715	T/C	0.71	-0.045(0.004)	9.63E-25
Creatine	Poultry	Food	0.083(0.013)	3.17E-11	CPS1	rs715	T/C	0.71	-0.045(0.004)	9.63E-25
Creatine	Traditional English	Pattern	0.076(0.013)	1.19E-09	CPS1	rs715	T/C	0.71	-0.045(0.004)	9.63E-25
Indolepropionate	Fruit & Vegetable	Pattern	0.082(0.009)	8.42E-21	ACSM2A	rs1394678	T/C	0.28	-0.035(0.004)	1.70E-20
Indolepropionate	Mediterranean Diet Score	Pattern	0.078(0.009)	1.60E-17	ACSM2A	rs1394678	T/C	0.28	-0.035(0.004)	1.70E-20
Indolepropionate	Bananas	Food	0.034(0.005)	1.05E-11	ACSM2A	rs1394678	T/C	0.28	-0.035(0.004)	1.70E-20
Indolepropionate	Apples & pears	Food	0.026(0.004)	2.39E-09	ACSM2A	rs1394678	T/C	0.28	-0.035(0.004)	1.70E-20
N-acetylornithine	Fruit & Vegetable	Pattern	0.050(0.009)	2.57E-08	NAT8	rs10206899	T/C	0.78	0.221(0.005)	4.66E-481
proline	Fruit & Vegetable	Pattern	-0.054(0.010)	1.95E-08	PRODH	rs2540641	A/C	0.08	0.063(0.004)	2.98E-59

Table 10. SNP associations with diet-associated metabolites from previous genome-wide association study (Shin et al., Nat Genet. 2014 Jun;46(6):543-50)

			Diet variable-M (Meta-analys	Metabolite is result)	<u></u>				SNP-Meta (Shin et al.	bolite , 2014)
Metabolite	Diet variable	Diet pattern or food	Beta(SE)	P	Gene symbol	SNP	EA/OA	EAF	Beta(SE)	Р
Pyroglutamine*	Low Meat	Pattern	0.120(0.013)	8.99E-20	SLC6A13	rs11613331	A/G	0.55	0.037(0.004)	2.23E-25
Pyroglutamine*	Low Meat	Pattern	0.120(0.013)	8.99E-20	SLC6A20	rs17279437	A/G	0.10	0.059(0.006)	1.25E-20
Pyroglutamine*	Poultry	Food	-0.091(0.013)	1.36E-12	SLC6A13	rs11613331	A/G	0.55	0.037(0.004)	2.23E-25
Pyroglutamine*	Poultry	Food	-0.091(0.013)	1.36E-12	SLC6A20	rs17279437	A/G	0.10	0.059(0.006)	1.25E-20
Pyroglutamine*	Other seafood	Food	-0.099(0.014)	5.22E-12	SLC6A13	rs11613331	A/G	0.55	0.037(0.004)	2.23E-25
Pyroglutamine*	Other seafood	Food	-0.099(0.014)	5.22E-12	SLC6A20	rs17279437	A/G	0.10	0.059(0.006)	1.25E-20
Pyroglutamine*	Meat	Food	-0.062(0.010)	2.10E-10	SLC6A13	rs11613331	A/G	0.55	0.037(0.004)	2.23E-25
Pyroglutamine*	Meat	Food	-0.062(0.010)	2.10E-10	SLC6A20	rs17279437	A/G	0.10	0.059(0.006)	1.25E-20
Tryptophan betaine	Fruit & Vegetable	Pattern	0.080(0.009)	1.94E-17	SLC22A4	rs2405522	A/G	0.17	-0.126(0.011)	4.49E-29
Tryptophan betaine	High Alcohol	Pattern	0.116(0.014)	1.08E-16	SLC22A4	rs2405522	A/G	0.17	-0.126(0.011)	4.49E-29
Tryptophan betaine	Mediterranean Diet Score	Pattern	0.084(0.011)	1.65E-15	SLC22A4	rs2405522	A/G	0.17	-0.126(0.011)	4.49E-29
Tryptophan betaine	Allium vegetables	Food	0.034(0.006)	4.73E-09	SLC22A4	rs2405522	A/G	0.17	-0.126(0.011)	4.49E-29
1,5-anhydroglucitol (1,5-AG)	Fruit & Vegetable	Pattern	-0.071(0.009)	1.02E-16	RAB3GA P1	rs7570971	A/C	0.33	-0.037(0.003)	7.86E-45
1,5-Anhydroglucitol (1,5-AG)	Fruit & Vegetable	Pattern	-0.071(0.009)	1.02E-16	MGAM	rs3800993	T/C	0.17	0.029(0.004)	1.53E-15
1,5-Anhydroglucitol (1,5-AG)	Other seafood	Food	-0.081(0.014)	1.80E-08	RAB3GA P1	rs7570971	A/C	0.33	-0.037(0.003)	7.86E-45
1,5-Anhydroglucitol (1,5-AG)	Other seafood	Food	-0.081(0.014)	1.80E-08	MGAM	rs3800993	T/C	0.17	0.029(0.004)	1.53E-15
1- Arachidonoylglyceroph osphoethanolamine*	Oily fish	Food	-0.097(0.015)	9.46E-11	FADS1	rs174578	A/T	0.66	0.056(0.003)	1.86E-94
1- Arachidonoylglyceroph osphoethanolamine*	Oily fish	Food	-0.097(0.015)	9.46E-11	SLCO1B 1	rs4149056	T/C	0.84	-0.040(0.003)	3.66E-31
1- Eicosatrienoylglycerop hosphocholine*	Oily fish	Food	-0.095(0.014)	1.12E-11	FADS1	rs968567	T/C	0.18	0.040(0.005)	2.84E-19
1- Eicosatrienoylglycerop hosphocholine*	Other seafood	Food	-0.082(0.014)	8.44E-09	FADS1	rs968567	T/C	0.18	0.040(0.005)	2.84E-19
1- Linoleoylglycerophosp hoethanolamine*	Oily fish	Food	-0.089(0.015)	1.60E-09	FADS1	rs174535	T/C	0.67	-0.044(0.004)	2.82E-36
10-Undecenoate (11:1n1)	Butter	Food	0.020(0.003)	2.35E-11	CYP4A11	rs9333029	A/G	0.87	-0.070(0.004)	1.52E-61

		_	Diet variable-M (Meta-analysi	/letabolite is result)	-				SNP-Metal (Shin et al.,	bolite 2014)
		Diet pattern			Gene					
Metabolite	Diet variable	or food	Beta(SE)	Р	symbol	SNP	EA/OA	EAF	Beta(SE)	Р
10-Undecenoate (11:1n1)	High Alcohol	Pattern	0.073(0.013)	8.17E-09	CYP4A11	rs9333029	A/G	0.87	-0.070(0.004)	1.52E-61
4-Androsten- 3beta,17beta-diol disulfate 1*	Wine	Food	0.041(0.003)	1.50E-31	SULT2A1	rs296396	T/C	0.17	-0.175(0.009)	1.48E-92
4-Androsten- 3beta,17beta-diol disulfate 1*	High Alcohol	Pattern	0.122(0.014)	3.27E-19	SULT2A1	rs296396	T/C	0.17	-0.175(0.009)	1.48E-92
4-Androsten- 3beta,17beta-diol disulfate 1*	Spirits and liquors	Food	0.040(0.007)	4.30E-09	SULT2A1	rs296396	T/C	0.17	-0.175(0.009)	1.48E-92
5-Alpha-androstan- 3beta,17beta-diol disulfate	Wine	Food	0.036(0.004)	2.71E-24	CYP3A5	rs10278040	A/G	0.04	-0.190(0.017)	1.17E-29
5-Alpha-androstan- 3beta,17beta-diol disulfate	Wine	Food	0.036(0.004)	2.71E-24	SULT2A1	rs2547231	A/C	0.83	0.081(0.008)	3.35E-22
5-Alpha-androstan- 3beta,17beta-diol disulfate	Wine	Food	0.036(0.004)	2.71E-24	ZCWPW1	rs13222543	T/C	0.02	-0.263(0.028)	1.18E-20
5-Alpha-androstan- 3beta,17beta-diol disulfate	High Alcohol	Pattern	0.112(0.014)	1.73E-15	CYP3A5	rs10278040	A/G	0.04	-0.190(0.017)	1.17E-29
5-Alpha-androstan- 3beta,17beta-diol disulfate	High Alcohol	Pattern	0.112(0.014)	1.73E-15	SULT2A1	rs2547231	A/C	0.83	0.081(0.008)	3.35E-22
5-Alpha-androstan- 3beta,17beta-diol disulfate	High Alcohol	Pattern	0.112(0.014)	1.73E-15	ZCWPW1	rs13222543	T/C	0.02	-0.263(0.028)	1.18E-20
Docosapentaenoate (n3 DPA; 22:5n3)	Wine	Food	0.024(0.003)	5.28E-15	FADS1	rs174538	A/G	0.30	-0.026(0.004)	8.90E-14
Docosapentaenoate (n3 DPA; 22:5n3)	Other seafood	Food	0.089(0.013)	3.44E-11	FADS1	rs174538	A/G	0.30	-0.026(0.004)	8.90E-14
Docosapentaenoate (n3 DPA; 22:5n3)	Oily fish	Food	0.080(0.013)	2.22E-10	FADS1	rs174538	A/G	0.30	-0.026(0.004)	8.90E-14
Eicosapentaenoate (EPA; 20:5n3)	Oily fish	Food	0.169(0.014)	1.57E-33	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22
Eicosapentaenoate (EPA; 20:5n3)	Fruit & Vegetable	Pattern	0.095(0.009)	1.86E-25	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22
Eicosapentaenoate (EPA; 20:5n3)	Other seafood	Food	0.154(0.015)	2.66E-24	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22
Eicosapentaenoate (EPA; 20:5n3)	Wine	Food	0.027(0.003)	6.63E-22	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22

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			Diet variable-M	/letabolite					SNP-Metal	bolite
		_	(Meta-analysi	is result)	-				(Shin et al.,	2014)
		Diet			Gono					
Metabolite	Diet variable	or food	Beta(SE)	Р	symbol	SNP	EA/OA	EAF	Beta(SE)	Р
Eicosapentaenoate (EPA: 20:5n3)	Mediterranean Diet Score	Pattern	0.083(0.009)	9.54E-21	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22
Eicosapentaenoate (EPA: 20:5n3)	Low Meat	Pattern	-0.123(0.014)	8.44E-19	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22
Eicosapentaenoate (EPA: 20:5n3)	Baked goods	Food	-0.014(0.002)	1.08E-12	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22
Eicosapentaenoate (EPA: 20:5n3)	Avocado	Food	0.108(0.018)	2.01E-09	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22
Eicosapentaenoate (EPA: 20:5n3)	High fat salad dressings	Food	0.044(0.008)	5.48E-08	FADS1	rs174556	T/C	0.30	-0.036(0.004)	1.97E-22
Epiandrosterone	Wine	Food	0.018(0.003)	6.29E-09	CYP3A5	rs11974702	A/G	0.91	0.184(0.010)	2.80E-75
Epiandrosterone	Wine	Food	0.018(0.003)	6.29E-09	ZCWPW1	rs13222543	T/C	0.02	-0.347(0.024)	3.31E-47
Myo-inositol	Wine	Food	0.022(0.003)	1.36E-15	SLC5A11	rs4788439	T/C	0.08	-0.027(0.004)	6.59E-13
Myo-inositol	Fruit & Vegetable	Pattern	0.052(0.008)	4.43E-11	SLC5A11	rs4788439	T/C	0.08	-0.027(0.004)	6.59E-13
Nonanoylcarnitine*	Butter	Food	0.026(0.003)	6.48E-17	ACADL	rs3738934	T/C	0.62	-0.106(0.004)	1.21E-134
Nonanoylcarnitine*	Butter	Food	0.026(0.003)	6.48E-17	THEM4	rs12566232	A/C	0.71	-0.045(0.005)	5.69E-19
Scyllo-inositol	Wine	Food	0.052(0.003)	1.47E-49	SLC5A11	rs4787294	A/T	0.93	0.075(0.008)	9.64E-21
Scyllo-inositol	High Alcohol	Pattern	0.143(0.014)	4.20E-26	SLC5A11	rs4787294	A/T	0.93	0.075(0.008)	9.64E-21
Scyllo-inositol	Fruit & Vegetable	Pattern	0.088(0.009)	2.22E-23	SLC5A11	rs4787294	A/T	0.93	0.075(0.008)	9.64E-21
Scyllo-inositol	Baked goods	Food	-0.017(0.003)	1.44E-10	SLC5A11	rs4787294	A/T	0.93	0.075(0.008)	9.64E-21
Scyllo-inositol	Mediterranean Diet Score	Pattern	0.066(0.010)	2.25E-10	SLC5A11	rs4787294	A/T	0.93	0.075(0.008)	9.64E-21
Scyllo-inositol	Fried fish	Food	-0.154(0.026)	4.44E-09	SLC5A11	rs4787294	A/T	0.93	0.075(0.008)	9.64E-21
Stearidonate (18:4n3)	Wine	Food	0.019(0.003)	3.17E-10	FADS1	rs174601	T/C	0.34	-0.034(0.004)	7.93E-16
1-Methylxanthine	Coffee	Food	0.010(0.002)	3.31E-09	NAT2	rs4921914	T/C	0.78	0.088(0.005)	1.09E-60
X-01911	High Alcohol	Pattern	0.115(0.014)	3.77E-17	COMT	rs4680	A/G	0.51	-0.044(0.006)	1.31E-13
X-01911	Wine	Food	0.026(0.003)	2.55E-14	COMT	rs4680	A/G	0.51	-0.044(0.006)	1.31E-13
X-02249	Butter	Food	0.026(0.003)	2.31E-16	CYP2C8	rs1934955	A/G	0.71	-0.030(0.004)	3.33E-16
X-02249	Low Meat	Pattern	-0.110(0.015)	7.70E-13	CYP2C8	rs1934955	A/G	0.71	-0.030(0.004)	3.33E-16
X-02249	Dieting	Pattern	0.091(0.014)	4.01E-11	CYP2C8	rs1934955	A/G	0.71	-0.030(0.004)	3.33E-16
X-02269	Oily fish	Food	0.175(0.013)	1.44E-38	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14
X-02269	Fruit & Vegetable	Pattern	0.102(0.009)	7.46E-31	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14

			Diet variable-M (Meta-analysi	/letabolite is result)					SNP-Meta (Shin et al.	bolite 2014)
		Diet pattern			Gene					
Metabolite	Diet variable	or food	Beta(SE)	Ρ	symbol	SNP	EA/OA	EAF	Beta(SE)	Р
X-02269	Other fish and seafood	Food	0.174(0.015)	9.25E-31	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14
X-02269	Mediterranean Diet Score	Pattern	0.099(0.009)	1.70E-29	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14
X-02269	Avocado	Food	0.147(0.019)	6.33E-15	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14
X-02269	Green leafy vegetables	Food	0.030(0.004)	2.39E-13	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14
X-02269	Low Meat	Pattern	-0.109(0.015)	2.61E-13	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14
X-02269	High fibre breakfast cereals	Food	0.030(0.004)	3.63E-12	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14
X-02269	Allium vegetables	Food	0.023(0.004)	3.55E-08	CYP2C8	rs2071426	T/C	0.71	0.055(0.007)	6.87E-14
X-08402	Butter	Food	0.017(0.003)	7.17E-10	SGPP1	rs7157785	T/G	0.16	0.068(0.003)	7.23E-87
X-08402	Butter	Food	0.017(0.003)	7.17E-10	SPTLC3	rs4814176	T/C	0.39	0.022(0.003)	2.97E-17
X-09789	Porridge	Food	0.094(0.008)	4.96E-33	SLC51A	rs7642243	C/G	0.39	0.048(0.006)	7.5E-16
X-09789	High Alcohol	Pattern	-0.116(0.013)	4.99E-19	SLC51A	rs7642243	C/G	0.39	0.048(0.006)	7.50E-16
X-09789	High fibre breakfast cereals	Food	0.038(0.005)	7.28E-14	SLC51A	rs7642243	C/G	0.39	0.048(0.006)	7.5E-16
X-09789	Fruit & Vegetable	Pattern	0.061(0.009)	3.50E-11	SLC51A	rs7642243	C/G	0.39	0.048(0.006)	7.50E-16
X-09789	Apples & pears	Food	0.020(0.004)	2.22E-08	SLC51A	rs7642243	C/G	0.39	0.048(0.006)	7.5E-16
X-09789	Wholemeal bread, grains	Food	0.015(0.003)	6.03E-08	SLC51A	rs7642243	C/G	0.39	0.048(0.006)	7.5E-16
X-10395	Wine	Food	0.031(0.003)	3.45E-26	NR1I3	rs4073054	A/C	0.63	0.016(0.002)	4.42E-19
X-10395	High Alcohol	Pattern	0.091(0.013)	3.80E-13	NR1I3	rs4073054	A/C	0.63	0.016(0.002)	4.42E-19
X-10510	Butter	Food	0.022(0.003)	8.28E-15	SGPP1	rs7157785	T/G	0.86	-0.044(0.004)	3.41E-37
X-11261	Low Meat	Pattern	0.105(0.014)	4.20E-14	SLC22A1	rs662138	C/G	0.83	0.063(0.006)	9.38E-25
X-11261	Low Meat	Pattern	0.105(0.014)	4.20E-14	SLC16A9	rs1171614	T/C	0.22	-0.042(0.006)	6.68E-14
X-11261	Fruit & Vegetable	Pattern	-0.061(0.010)	2.08E-10	SLC22A1	rs662138	C/G	0.83	0.063(0.006)	9.38E-25
X-11261	Fruit & Vegetable	Pattern	-0.061(0.010)	2.08E-10	SLC16A9	rs1171614	T/C	0.22	-0.042(0.006)	6.68E-14
X-11315	Fruit & Vegetable	Pattern	0.142(0.008)	3.87E-63	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Mediterranean Diet Score	Pattern	0.109(0.009)	7.94E-34	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Nuts	Food	0.054(0.005)	3.75E-25	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Apples & pears	Food	0.035(0.004)	9.63E-20	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12

	Table 10. SNP associations with diet-associated metabolites from	previous genome-wide association study (Shin et al., Nat Genet. 2014 Jun;46(6):543-50)
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			Diet variable-M (Meta-analysi	letabolite is result)	_				SNP-Meta (Shin et al.	bolite , 2014)
Metabolite	Diet variable	Diet pattern or food	Beta(SE)	Р	Gene symbol	SNP	EA/OA	EAF	Beta(SE)	Р
X-11315	Oily fish	Food	0.106(0.014)	1.62E-14	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Berries	Food	0.103(0.013)	2.70E-14	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Peaches	Food	0.127(0.017)	5.92E-14	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Green leafy vegetables	Food	0.033(0.005)	1.71E-12	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	White and brown bread, refined grains	Food	-0.017(0.002)	1.84E-12	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Confectionary & jams	Food	-0.010(0.001)	1.53E-11	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Traditional English	Pattern	-0.093(0.014)	1.68E-11	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Savoury pies	Food	-0.221(0.034)	9.08E-11	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Citrus fruit	Food	0.026(0.004)	6.50E-10	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Fried fish	Food	-0.164(0.027)	1.54E-09	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	High fibre breakfast cereals	Food	0.030(0.005)	2.75E-09	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11315	Baked goods	Food	-0.012(0.002)	2.90E-08	SLC6A20	rs4327428	A/C	0.11	-0.031(0.004)	2.81E-12
X-11381	Mediterranean Diet Score	Pattern	-0.069(0.009)	8.27E-14	SLC16A9	rs12356193	A/G	0.84	0.024(0.003)	8.36E-20
X-11381	High Alcohol	Pattern	-0.077(0.012)	4.54E-11	SLC16A9	rs12356193	A/G	0.84	0.024(0.003)	8.36E-20
X-11381	Processed meats	Food	0.049(0.008)	4.74E-10	SLC16A9	rs12356193	A/G	0.84	0.024(0.003)	8.36E-20
X-11381	Traditional English	Pattern	0.072(0.012)	3.71E-09	SLC16A9	rs12356193	A/G	0.84	0.024(0.003)	8.36E-20
X-11381	Soy foods	Food	-0.108(0.020)	5.80E-08	SLC16A9	rs12356193	A/G	0.84	0.024(0.003)	8.36E-20
X-11469	Oily fish	Food	0.176(0.013)	5.87E-39	CYP2C8	rs2071426	T/C	0.71	0.052(0.007)	3.97E-14
X-11469	Oily fish	Food	0.176(0.013)	5.87E-39	SLC17A3	rs11754288	A/G	0.44	0.043(0.006)	1.88E-12
X-11469	Fruit & Vegetable	Pattern	0.106(0.009)	3.39E-34	CYP2C8	rs2071426	T/C	0.71	0.052(0.007)	3.97E-14
X-11469	Other fish and seafood	Food	0.175(0.015)	2.99E-31	CYP2C8	rs2071426	T/C	0.71	0.052(0.007)	3.97E-14
X-11469	Other fish and seafood	Food	0.175(0.015)	2.99E-31	SLC17A3	rs11754288	A/G	0.44	0.043(0.006)	1.88E-12
X-11469	Mediterranean Diet Score	Pattern	0.100(0.009)	9.16E-31	CYP2C8	rs2071426	T/C	0.71	0.052(0.007)	3.97E-14
X-11469	Avocado	Food	0.153(0.019)	9.36E-16	CYP2C8	rs2071426	T/C	0.71	0.052(0.007)	3.97E-14
X-11469	Avocado	Food	0.153(0.019)	9.36E-16	SLC17A3	rs11754288	A/G	0.44	0.043(0.006)	1.88E-12

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			Diet variable-N (Meta-analysi	letabolite s result)				_	SNP-Metabolite (Shin et al., 2014)		
Metabolite	Diet variable	Diet pattern or food	Bota(SE)	P	Gene	SNP	EA/OA	FAF	Bota(SE)	Þ	
X-11469	Green leafy	Food	0.030(0.004)	5.66E-14	CYP2C8	rs2071426	T/C	0.71	0.052(0.007)	3.97E-14	
	vegetables								, , ,		
X-11469	Green leafy vegetables	Food	0.030(0.004)	5.66E-14	SLC17A3	rs11754288	A/G	0.44	0.043(0.006)	1.88E-12	
X-11469	Low Meat	Pattern	-0.110(0.015)	1.88E-13	SLC17A3	rs11754288	A/G	0.44	0.043(0.006)	1.88E-12	
X-11469	High fibre breakfast cereals	Food	0.031(0.004)	5.40E-13	CYP2C8	rs2071426	T/C	0.71	0.052(0.007)	3.97E-14	
X-11469	High fibre breakfast cereals	Food	0.031(0.004)	5.40E-13	SLC17A3	rs11754288	A/G	0.44	0.043(0.006)	1.88E-12	
X-11478	Low Meat	Pattern	0.094(0.015)	9.86E-11	ACSM2A	rs6497490	T/G	0.88	-0.082(0.008)	4.95E-27	
X-11550	Wine	Food	0.018(0.003)	3.94E-11	CETP	rs247616	T/C	0.32	0.010(0.001)	1.11E-11	
X-11799	Wine	Food	0.033(0.003)	1.66E-27	GBA3	rs3099557	A/G	0.85	-0.148(0.011)	1.83E-39	
X-11799	High Alcohol	Pattern	0.120(0.014)	6.07E-18	GBA3	rs3099557	A/G	0.85	-0.148(0.011)	1.83E-39	
X-11799	Baked goods	Food	-0.017(0.003)	4.88E-09	GBA3	rs3099557	A/G	0.85	-0.148(0.011)	1.83E-39	
X-12038	Wine	Food	0.022(0.003)	9.20E-13	CETP	rs1800775	A/C	0.50	0.013(0.002)	1.02E-11	
X-12038	High Alcohol	Pattern	0.064(0.012)	1.02E-07	CETP	rs1800775	A/C	0.50	0.013(0.002)	1.02E-11	
X-12063	Fruit & Vegetable	Pattern	-0.048(0.008)	2.31E-09	СҮРЗА5	rs10242455	A/G	0.93	0.221(0.010)	1.67E-109	
X-12063	Fruit & Vegetable	Pattern	-0.048(0.008)	2.31E-09	SLCO1B 1	rs4149056	T/C	0.84	-0.118(0.007)	1.26E-73	
X-12627	Wine	Food	0.022(0.003)	5.88E-15	ELOVL2	rs4713169	C/G	0.42	-0.032(0.004)	2.87E-14	
X-12627	Oily fish	Food	0.073(0.012)	5.74E-09	ELOVL2	rs4713169	C/G	0.42	-0.032(0.004)	2.87E-14	
X-12798	Low fat milk	Food	0.062(0.008)	1.24E-15	SLC22A1	rs316019	A/C	0.09	-0.181(0.005)	1.56E-259	
X-12798	Low fat milk	Food	0.062(0.008)	1.24E-15	SLC16A9	rs1171615	T/C	0.78	-0.048(0.004)	3.19E-34	
X-12798	High Alcohol	Pattern	-0.099(0.012)	1.58E-15	SLC22A1	rs316019	A/C	0.09	-0.181(0.005)	1.56E-259	
X-12798	Mediterranean Diet Score	Pattern	-0.069(0.010)	6.80E-13	SLC16A9	rs1171615	T/C	0.78	-0.048(0.004)	3.19E-34	
X-13477	Mediterranean Diet Score	Pattern	0.067(0.010)	1.56E-10	NAT8	rs10206899	T/C	0.78	0.025(0.003)	1.73E-14	
X-14473	Coffee	Food	0.038(0.001)	6.12E- 187	CYP2C9	rs4986894	T/C	0.84	-0.042(0.006)	5.55E-12	
X-14473	Black tea	Food	-0.024(0.001)	1.36E-72	CYP2C9	rs4986894	T/C	0.84	-0.042(0.006)	5.55E-12	
X-14473	Dieting	Pattern	-0.148(0.014)	8.46E-25	CYP2C9	rs4986894	T/C	0.84	-0.042(0.006)	5.55E-12	
X-21365 [trimethyl-N- aminovalerate]	Low fat milk	Food	0.076(0.007)	9.36E-27	SLC22A4	rs273913	T/C	0.38	0.026(0.003)	1.08E-25	

Table 10. SNP associations with diet-associated metabolites from previous genome-wide association study (Shin et al., Nat Genet. 2014 Jun;46(6):543-50)

			Diet variable-Metabolite (Meta-analysis result)						SNP-Metal (Shin et al.,	bolite , 2014)
Metabolite	Diet variable	Diet pattern or food	Beta(SE)	Р	Gene symbol	SNP	EA/OA	EAF	Beta(SE)	P
X-21365 [trimethyl-N- aminovalerate]	Low fat milk	Food	0.076(0.007)	9.36E-27	MARCH8	rs2291429	A/C	0.76	0.017(0.003)	8.69E-11
X-21365 [trimethyl-N- aminovalerate]	Mediterranean Diet Score	Pattern	-0.073(0.009)	1.48E-15	SLC22A4	rs273913	T/C	0.38	0.026(0.003)	1.08E-25
X-21365 [trimethyl-N- aminovalerate]	Mediterranean Diet Score	Pattern	-0.073(0.009)	1.48E-15	MARCH8	rs2291429	A/C	0.76	0.017(0.003)	8.69E-11
X-21365 [trimethyl-N- aminovalerate]	High Alcohol	Pattern	-0.091(0.012)	3.91E-15	SLC22A4	rs273913	T/C	0.38	0.026(0.003)	1.08E-25
X-21365 [trimethyl-N- aminovalerate]	High Alcohol	Pattern	-0.091(0.012)	3.91E-15	MARCH8	rs2291429	A/C	0.76	0.017(0.003)	8.69E-11

Table 10. SNP associations with diet-associated metabolites from previous genome-wide association study (Shin et al., Nat Genet. 2014 Jun;46(6):543-50)

Notes: Statistics are reported for the most associated metabolite at each single nucleotide polymorphism (SNP) for those metabolites associated with food intakes and diet patterns. The genome-wide association study was previously performed on the TwinsUK and KORA blood metabolite datasets (Shin et al., Nat Genet. 2014 Jun;46(6):543-50). EA/OA = effect/other allele; EAF = effect allele frequency.

			Food Group-Metabolite (Meta-analysis result)		_	SNP-Metabolite (Shin et al., 2014)		Food variable-SNP			
Metabolite	Diet variable	Diet pattern or food	Beta(SE)	P	SNP	Beta(SE)	P	Beta(SE)	Р	Gene symbol	
3-Phenylpropionate (hydrocinnamate)	Fruit & Vegetable	Pattern	0.080(0.010)	1.73E-16	rs11647589	0.030(0.005)	2.85E-11	0.140(0.059)	1.80E-02	ACSM5	
3-Phenylpropionate (hydrocinnamate)	Savoury pies	Food	-0.231(0.037)	3.72E-10	rs11647589	0.030(0.005)	2.85E-11	-0.023(0.012)	4.76E-02	ACSM5	
Indolepropionate	Fruid & Vegetable	Pattern	0.082(0.009)	8.42E-21	rs1394678	-0.035(0.004)	1.70E-20	0.163(0.065)	1.20E-02	ACSM2A	
Indolepropionate	Mediterranean Diet Score	Pattern	0.078(0.009)	1.60E-17	rs1394678	-0.035(0.004)	1.70E-20	0.125(0.053)	1.80E-02	ACSM2A	
Pyroglutamine*	Meat	Food	-0.062(0.010)	2.10E-10	rs17279437	0.059(0.006)	1.25E-20	-0.142(0.069)	3.95E-02	SLC6A20	
1,5-Anhydroglucitol (1,5-AG)	Other seafood	Food	-0.081(0.014)	1.80E-08	rs3800993	0.029(0.004)	1.53E-15	-0.090(0.034)	8.51E-03	MGAM	
5-Alpha-androstan- 3beta,17beta-diol disulfate	High Alcohol	Pattern	0.112(0.014)	1.73E-15	rs10278040	-0.190(0.017)	1.17E-29	0.198(0.077)	1.10E-02	CYP3A5	
5-Alpha-androstan- 3beta,17beta-diol disulfate	High Alcohol	Pattern	0.112(0.014)	1.73E-15	rs13222543	-0.263(0.028)	1.18E-20	0.301(0.129)	2.00E-02	ZCWPW1	
Myo-inositol	Wine	Food	0.022(0.003)	1.36E-15	rs4788439	-0.027(0.004)	6.59E-13	-0.654(0.236)	5.59E-03	SLC5A11	
Scyllo-inositol	Fried fish	Food	-0.154(0.026)	4.44E-09	rs4787294	0.075(0.008)	9.64E-21	0.088(0.035)	1.25E-02	SLC5A11	
Scyllo-inositol	Wine	Food	0.052(0.003)	1.47E-49	rs4787294	0.075(0.008)	9.64E-21	-0.586(0.243)	1.60E-02	SLC5A11	
X-02269	Green leafy vegetables	Food	0.030(0.004)	2.39E-13	rs2071426	0.055(0.007)	6.87E-14	0.228(0.110)	3.76E-02	CYP2C8	
X-08402	Butter	Food	0.017(0.003)	7.17E-10	rs7157785	0.068(0.003)	7.23E-87	0.004(0.001)	2.95E-03	SGPP1	
X-10395	High Alcohol	Pattern	0.091(0.013)	3.80E-13	rs4073054	0.016(0.002)	4.42E-19	-0.071(0.034)	4.10E-02	NR1/3	
X-10510	Butter	Food	0.022(0.003)	8.28E-15	rs7157785	-0.044(0.004)	3.41E-37	0.004(0.001)	2.95E-03	SGPP1	
X-11315	Citrus fruit	Food	0.026(0.004)	6.50E-10	rs4327428	-0.031(0.004)	2.81E-12	0.473(0.209)	2.37E-02	SLC6A20	
X-11315	Traditional English	Pattern	-0.093(0.014)	1.68E-11	rs4327428	-0.031(0.004)	2.81E-12	-0.114(0.057)	4.80E-02	SLC6A20	
X-11469	Green leafy vegetables	Food	0.030(0.004)	5.66E-14	rs2071426	0.052(0.007)	3.97E-14	0.228(0.110)	3.76E-02	CYP2C8	
X-11469	Avocado	Food	0.153(0.019)	9.36E-16	rs11754288	0.043(0.006)	1.88E-12	0.043(0.022)	4.75E-02	SLC17A3	

Table 11. Diet-SNP associations with SNPs associated with metabolites from a previous genome-wide association study (Shin et al., Nat Genet. 2014 Jun;46(6):543-50)

Notes: Statistics are reported for associations between food intake at metabolite-SNPs which meet the 5% level of significance. A linear regression was performed using the applicable metabolite-associated SNP as a predictor of reported food group intake or dietary pattern score adjusted for age, energy intake and family relatedness; an additive genetic model was performed for each. No associations met statistical significance of 4.76x10⁻⁴ (0.05/105 tests). The genome-wide association study was previously performed on the TwinsUK and KORA blood metabolite datasets (Shin et al., Nat Genet. 2014 Jun;46(6):543-50).

Appendix D. Chapter 5 Appendices

Document 1: Food Preference Questionnaire

Your food and lifestyle preferences

Welcome to our latest questionnaire.

Throughout this section of the questionnaire you will use the scales shown below to report your likes and dislikes for foods, physical activities, and experiences.

The term 'Dislike' ranges from weak disliking (close to neutral) to strongest disliking of any kind (left-hand side of the scale).

The term 'Like' is right of neutral, ranging from weaker liking (close to neutral) to strongest liking of any kind (right-hand side of the scale). The term **'Neutral**' is in the middle (neither like nor dislike; "it's okay").

Questions marked with (*) are mandatory.

Below are some examples to show you how to use the scales.

The following two examples are how a person would rate experiences that they dislike:





The following example is how a person would rate an experience that they neither like nor dislike (a neutral experience):



Page 1

* Please rate your liking or disliking of the following.

The ends of the scale are the strongest disliking or liking of any kind. You can place your rating anywhere between the labels.

To place your rating on the scale, either click the position on the scale that you would like to choose, or drag the slider to that place. If your rating is neutral, simply click on the slider in its starting position at the center of the scale. If you have never tried a food, physical activity or experience, then select the "never tried" option.

No.	neither like nor dislike	strongest liking of
By Theor of any kind	"it's okay"	any kind
headlights Hearing your favourite piece of music		

	Never tree	strongest disliking	() () () () () () () () () () () () () (ither like nor dislike	strongest liking
26	broccoli 🔲			<u> </u>	
A. B	tortilla chips or crisps	I <u> </u>		I]
	crispy bacon	l]
	beer	1		I]
	prawns & 🔲 shellfish	۱ـــــــــــــــــــــــــــــــــــــ]
X	butter/ margarine 🔲	ا ل		ĺ]
	chips 🔲	ıL		Ĩ]
	wholemeal Development	I			
bl	ue cheese	IL		I	
	skimmed milk	ıL	Page 3	I]

Neve (Na)	tied	strongest disliking	()) nei	ither like nor dislike	٩	strongest liking
fresh tomatoes				ĺ		of any kind
black coffee	Q	L]
cooling off a hot d	on ay				a]
pizza		L		I		
plain yoghurt	Q	L]
frie chicke	n 🗖	L		<u>L</u>]
burn spicy foo	of ds	[]
bag rolls		L]
sausa	ge 🔲	L		I]
soy sau	ce 🔲	L		I]

Page 4

	Neve (NS)	tied	strongest disliking of any kind		either like nor dislike	٢	strongest liking
ڔٵٛ	king the stairs		L				
	fizzy so drinks & swee drinks	ft s t s(e.g.	lemonade, squash	es)	ĺ		
69	salty pretzels						
	fresh coriander		<u> </u>	,			
	pasta/ noodles		L		Ĩ		
""	going to the gym		L		1]
🌔 ma	ayonnaise		[]		L]
E	chili pepper						I
	whole milk]
	baked chicken				- 1]
				Page 5			

To place your rating on the scale, either click the position on the scale that you would like to choose, or drag the slider to that place. If your rating is neutral, simply click on the slider in its starting position at the center of the scale. If you have never tried a food, physical activity or experience, then select the "never tried" option.

	Never As	tried	strongest disliking of any kind	neither like nor dislike	٩	strongest liking of any kind
-	olives	Q	<u> </u>]
	dark chocolate		<u> </u>]
	garlic		L	ĺ]
Ø	ketchup			[
	ice cream		L	 [
	grapefruit		<u> </u>	 		
S	corn flakes			 1		
	exercisin alone	g	L	 I		
W	hite rice			 ſ]
	Smell of freshly cut grass	•		 ľ		J

To place your rating on the scale, either click the position on the scale that you would like to choose, or drag the slider to that place. If your rating is neutral, simply click on the slider in its starting position at the center of the scale. If you have never tried a food, physical activity or experience, then select the "never tried" option.

	Neve As	lied	strongest disliking of any kind	neither like n	or dislike	strongest liking of any kind
	salad dressing	Q		I.]
-0	fried fish	D				
J	banana	Q	1		8]
	pork chops	Q	L	 I		
	lentils/ beans		L]
	commutin (car, bus, train)	g	L	1]
5	red wine		<u>[</u>	 		
	pear		[1]
i i i i i i i i i i i i i i i i i i i	cake icing			I]
×	vhite potato	[,] D	<u> </u>	 Í		

To place your rating on the scale, either click the position on the scale that you would like to choose, or drag the slider to that place. If your rating is neutral, simply click on the slider in its starting position at the center of the scale. If you have never tried a food, physical activity or experience, then select the "never tried" option.

	Mayar High May High melon (yellow or green)	strongest disliking of any kind	eeither like nor dislike	())	strongest liking of any kind
m	saut éed ushrooms)[
0	playing sports	h	 ĺ		
	extra virgin blive oil				[
and the second	asparagus		I]
e W	vith others	۱ <u>ـــــ</u>	I]
	raw carrots		T]
5.00	going to pub <i>l</i> bar			-]
	jam/jelly	<u>ا</u> ـــــــ	 I	<u></u>]
	bicycling	<u>ا</u> لــــــــــ]

To place your rating on the scale, either click the position on the scale that you would like to choose, or drag the slider to that place. If your rating is neutral, simply click on the slider in its starting position at the center of the scale. If you have never tried a food, physical activity or experience, then select the "never tried" option.

HA CON	Never (NA)	tied	strongest disliking of any kind) (intermediate and intermediate and int) 🧿) (interpretation of any kind)
	white wine			 []
	watching television	Q		I]
	lemon		_	 ĺ]
	porridge		L]
	cherries		<u> </u>	 [
	vinegar	Q	L]
	coffee or tea with sug	ar	<u> </u>]
	beetroot		[ĺ]
	horseradish wasabi	` □	l	I		
	working up a sweat			ľ		I

To place your rating on the scale, either click the position on the scale that you would like to choose, or drag the slider to that place. If your rating is neutral, simply click on the slider in its starting position at the center of the scale. If you have never tried a food, physical activity or experience, then select the "never tried" option.

Novas	tried	strongest disliking	(i)) (i))	neither like nor disli	ke	strongest liking
vodka, gin, scotch						of any kind
tuna or salmon		Ĺ]
getting caught in a lie		L]
biscuits, cakes or pastries		<u> </u>	<u>e</u>	ĺ]
ham		L		ĺ]
high-fibre bar		<u> </u>		Í]
strawberries		[]
savoury biscuits		<u>L</u>		I]
gherkins]
seeing a mouse at home	t 🗖	L		1]

Never (NS)	tien strongest disliking	() Inel	ither like nor dislike	٩	strongest liking
spinach/ greens					
pineapple	° 🖸 L]
hot tea			ĺ]
cheese cake		75.]
salting food			Í]
char- grilled meats					
aubergine			<u> </u>		J
cheddar cheese	" • • • •]
Tabasco sauce	° Q [I]
raw onion		Page 11			
Use the scales below to report your likes and dislikes for foods, physical activities, and experiences.

To place your rating on the scale, either click the position on the scale that you would like to choose, or drag the slider to that place. If your rating is neutral, simply click on the slider in its starting position at the center of the scale. If you have never tried a food, physical activity or experience, then select the "never tried" option.

New Cast	a strongest disliking of any kind		ither like nor dislike	strongest liking
going to a café			Ī	
	^{ge}		I]
sweet coffee drinks & whipped cream			I]
black pepper			I]
cigarett smoking]
faux meat products (e.g. Quo	t I I I I I I I I I I I I I I I I I I I		1]
curries			1]
diet fizzy soff drinks (e.g. Coke Zo Diet Coke)	ero,		I]
unsalted nuts			I	
beef steak	, D	Page 12	I]

Use the scales below to report your likes and dislikes for foods, physical activities, and experiences.

To place your rating on the scale, either click the position on the scale that you would like to choose, or drag the slider to that place. If your rating is neutral, simply click on the slider in its starting position at the center of the scale. If you have never tried a food, physical activity or experience, then select the "never tried" option.

No. By Tray, Strongest disliking of any kind	neither like nor dislike	strongest liking of any kind
eggs 🔲	I]
soy milk	I]
low calorie sweetener in coffee, teas or other beverages (e.g. Canderel)	I]

Food avoidance questions:

Α	۱						В				
Do you avoid items listed	dany o below 1	of the ?	lf yes, w	hatis/aret	he reason(s)	why you avoid	this food?	? (select all tha	t apply)		
			(1) I am allergic	(2) I am intolerant (e.g. Impaired digestion of the food)	(3) I am sensitive (e.g. I feel discomfort following eating)	(4) I have an illness/ disease (e.g. Inflammatory bowel disease)	(5) I believe it is better for long- term health (e.g. reduced risk of chronic disease)	(6) I believe it is better for weight loss/ maintenance	(7) Because of my beliefs (e.g. religion, veganism)	(8) I dislike this food	(9) Other
	(0) No	(1) Yes									
1. Fish	1	1		1	1	1	E.				L
2. Shellfish	3	I.		1	L.	1	I.				E.
3. Red meat	10	E.		٦	٦	٦	7	0.00	-8	10-00	٦
4. Poultry	Г	E.	-	٦	٦	٦	٦	8 	-8	-	٦
5. Gluten	Г	Г	la d	٦	٦	٦	٦	6	-20	1000	٦
6. Dairy	Г	Г		٦	Г	٦	٦	0.00		-	٦
7. Eggs	Г	Г	100	٦	٦	٦	٦	1 		1000	٦
8. Sweets	1 T	Γ.		٦	Г	٦	٦	6		1000	٦
9. Soy	Г	E.	=	٦	٦	٦	٦	1		1	٦
10. Nuts	Г	Г	100	٦	٦	٦	٦	8,00		1.00	٦
11. Alcohol] ⊺ ≊	Г		٦	٦	7	٦	0	-33	11 111	

2. Do you avoid any other foods?(0) No (skip to question 3)

(1) Yes, please complete the table below..

A					В				
What other foods do you avoid?	lf yes, w	/hatis/aret	he reason(s)	why you avoid	this food	? (select all tha	tapply)		
Please click on the box and type only one food type per box.	(1) I am allergic	(2) I am intolerant (e.g. Impaired digestion of the food)	(3) I am sensitive (e.g. I feel discomfort following eating)	(4) I have an illness/ disease (e.g. Inflammatory bowel disease)	(5) I believe it is better for long- term health (e.g. reduced risk of chronic disease)	(6) I believe it is better for weight loss/ maintenance	(7) Because of my beliefs (e.g. religion, veganism)	(8) I dislike this food	(9) Other
12. Other:		Ì	l I	. 1	1				I.
13. Other:		1	l I	1	L I				I.
14. Other:		1	i.	1	I.				ľ
15. Other:		Ĩ	i.	. 1	1				I.
16. Other:		1	L.	1					

Eating rate questions:

3. How would you cor	npare your eating	rate to oth	iers?			
(1) □ very slow	(2) [−] relative	ly slow	(3) [–] medium	(4) ⊓ relat	tively fast	(5) ⊓ very fast
4. Do you usually eat	until you feel full?	(2) = 1	quit eating when I fee	al full (3)		eating after I feel full
5. Between you and y	our twin: Who us	ually eats t	he most?		The continue of	
(1) [–] Me (2) ⊓ We eat abou	t the same	amount (3) ⊓ My twin	(99999	902) ⊓ Don't know

Press SUBMIT to complete your questionnaire, otherwise, press BACK button to modify your answers.

THANK YOU VERY MUCH FOR TAKING THE TIME TO COMPLETE THIS QUESTIONNAIRE. WE APPRECIATE ALL YOUR HELP

				Discov	very	Discorda	ant	_	Meta-an	alysis
Food		Super-								
group	Metabolite	pathway	Sub-pathway	Beta (SE)	Р	Beta (SE)	Р		Beta (SE)	Р
Vegetables	X-11315			0.009 (0.002)	1.06E-08	0.007 (0.005)	1.51E-01		0.009 (0.002)	2.94E-09
	X-11372			-0.007 (0.001)	1.09E-06	-0.004 (0.004)	3.06E-01		-0.007 (0.001)	6.20E-07
	X-12063			-0.006 (0.001)	2.36E-06	-0.002 (0.003)	4.28E-01		-0.005 (0.001)	2.33E-06
Fruit	stachydrine	Xenobiotics	Food component, Plant	0.021 (0.002)	3.65E-17	0.012 (0.004)	6.86E-03		0.019 (0.002)	8.71E-19
	glycerate	Carbohydrate	Glycolysis, gluconeogenesis, pyruvate metabolism	0.017 (0.002)	7.92E-13	0.017 (0.004)	2.33E-04		0.017 (0.002)	2.08E-16
	threonate	Cofactors and vitamins	Ascorbate and aldarate metabolism	0.012 (0.002)	2.47E-08	0.015 (0.004)	3.38E-04		0.013 (0.002)	1.89E-11
	threitol	Carbohydrate	Nucleotide sugars, pentose metabolism	0.012 (0.002)	5.18E-10	0.005 (0.003)	1.19E-01		0.010 (0.002)	7.27E-10
	docosahexaenoate (DHA; 22:6n3)	Lipid	Essential fatty acid	0.011 (0.002)	6.54E-06	0.010 (0.004)	2.13E-02		0.011 (0.002)	3.42E-07
	scyllo-inositol	Lipid	Inositol metabolism	0.012 (0.002)	1.03E-06	0.007 (0.004)	1.11E-01		0.011 (0.002)	3.68E-07
	proline	Amino acid	Urea cycle; arginine-, proline-, metabolism	-0.010 (0.002)	1.70E-06	-0.005 (0.005)	3.33E-01		-0.009 (0.002)	1.30E-06
	hippurate	Xenobiotics	Benzoate metabolism	0.011 (0.002)	7.86E-07	0.003 (0.004)	4.69E-01		0.009 (0.002)	2.12E-06
	X-11315			0.021 (0.002)	1.08E-16	0.014 (0.004)	1.43E-03		0.019 (0.002)	2.52E-19
	X-11372			-0.012 (0.002)	5.68E-08	-0.007 (0.004)	8.31E-02		-0.011 (0.002)	1.66E-08
Whole grains	3-phenylpropionate (hydrocinnamate)	Amino acid	Phenylalanine & tyrosine metabolism	0.017 (0.003)	1.90E-07	0.003 (0.007)	6.79E-01		0.015 (0.003)	7.90E-07
	eicosapentaenoate (EPA; 20:5n3)	Lipid	Essential fatty acid	0.015 (0.003)	2.30E-07	0.001 (0.006)	9.09E-01		0.013 (0.003)	1.67E-06
	X-09789			0.021 (0.003)	1.19E-12	0.015 (0.006)	1.00E-02		0.020 (0.003)	2.65E-14
	X-11372			-0.022 (0.003)	7.00E-13	-0.005 (0.006)	3.43E-01		-0.018 (0.003)	6.97E-12
	X-11469			0.019 (0.003)	4.72E-11	0.003 (0.007)	6.77E-01		0.017 (0.003)	2.64E-10
	X-02269			0.019 (0.003)	1.09E-10	0.001 (0.007)	8.54E-01		0.017 (0.003)	7.75E-10
	X-11315			0.015 (0.003)	3.95E-06	0.014 (0.005)	1.42E-02		0.014 (0.003)	1.37E-07
Nuts and legumes	tryptophan betaine	Amino acid	Tryptophan metabolism	0.048 (0.007)	9.22E-12	0.017 (0.011)	1.49E-01		0.040 (0.006)	2.52E-11
	X-11315			0.030 (0.006)	7.82E-07	0.023 (0.012)	6.24E-02		0.028 (0.005)	1.18E-07
Seafood	3-carboxy-4-methyl- 5-propyl-2- furanpropanoate (CMPF)	Lipid	Fatty acid, dicarboxylate	0.149 (0.014)	7.86E-24	0.112 (0.026)	2.84E-05		0.140 (0.012)	4.47E-29
	docosahexaenoate (DHA; 22:6n3)	Lipid	Essential fatty acid	0.167 (0.016)	5.32E-25	0.085 (0.022)	1.96E-04		0.139 (0.013)	1.03E-27
	eicosapentaenoate (EPA; 20:5n3)	Lipid	Essential fatty acid	0.133 (0.015)	4.36E-19	0.100 (0.027)	3.97E-04		0.126 (0.013)	1.25E-22
	1- docosahexaenoylglyc	Lipid	Lysolipid	0.103 (0.013)	4.42E-14	0.050 (0.030)	9.59E-02		0.094 (0.012)	1.47E-14

				Discov	very	_	Discorda	ant		Meta-an	alysis
Food group	Metabolite	Super- pathway	Sub-pathway	Beta (SE)	Р		Beta (SE)	Р		Beta (SE)	Р
5.01	ero-phosphocholine*										
	docosapentaenoate (n3 DPA; 22:5n3)	Lipid	Essential fatty acid	0.088 (0.014)	1.62E-10		0.038 (0.023)	1.06E-01		0.075 (0.012)	1.42E-10
	1- arachidonoylglycer	Lipid	Lysolipid	-0.068 (0.013)	4.51E-07		-0.117 (0.028)	7.99E-05		-0.077 (0.012)	1.99E-10
	o- phosphoethanolami ne*										
	pvroglutamine*	Amino acid	Glutamate metabolism	-0.072 (0.014)	7.09E-07		-0.079 (0.021)	2.52E-04		-0.074 (0.012)	3.54E-10
	1-oleoylglycero- phosphoethanolami ne	Lipid	Lysolipid	-0.076 (0.013)	2.99E-09		-0.052 (0.027)	5.66E-02		-0.072 (0.012)	4.28E-10
	pseudouridine	Nucleotide	Pyrimidine metabolism, uracil containing	-0.071 (0.013)	1.04E-07		-0.051 (0.018)	6.92E-03		-0.064 (0.011)	2.40E-09
	1- eicosatrienoylglyce ro-phosphocholine*	Lipid	Lysolipid	-0.062 (0.013)	2.28E-06		-0.092 (0.026)	7.74E-04		-0.068 (0.012)	6.45E-09
	1-linoleoylglycero- phosphoethanolami ne*	Lipid	Lysolipid	-0.064 (0.014)	4.08E-06		-0.058 (0.028)	4.26E-02		-0.063 (0.012)	4.01E-07
	X-11469			0.140 (0.014)	2.17E-21		0.120 (0.028)	4.19E-05		0.136 (0.013)	2.04E-26
	X-02269			0.139 (0.015)	2.27E-20		0.121 (0.027)	2.49E-05		0.135 (0.013)	1.45E-25
	X-11315			0.078 (0.014)	1.82E-08		0.044 (0.025)	7.47E-02		0.070 (0.012)	5.35E-09
	X-12627			0.076 (0.013)	1.61E-08		0.026 (0.023)	2.53E-01		0.064 (0.012)	3.62E-08
	X-12644			0.087 (0.015)	9.74E-09		0.008 (0.019)	6.84E-01		0.057 (0.012)	1.39E-06
	X-11437			0.076 (0.015)	3.97E-07		0.009 (0.027)	7.53E-01		0.061 (0.013)	3.41E-06
	X-12798			-0.061 (0.013)	6.13E-06		-0.035 (0.035)	3.18E-01		-0.058 (0.013)	4.15E-06
White meat	3-methylhistidine	Amino acid	Histidine metabolism	0.138 (0.023)	4.46E-09		0.136 (0.046)	3.97E-03		0.138 (0.021)	3.63E-11
Red,	trans-4-	Amino acid	Urea cycle; arginine-,	0.039 (0.007)	5.30E-08		0.014 (0.015)	3.57E-01		0.034 (0.006)	8.32E-08
processed most and	nydroxyproline	Amino opid	Creating metabolism	0.027 (0.007)			0.016 (0.014)	2.625.04		0.022 (0.006)	7 265 07
		Amino acio	Creatine metabolism	0.037(0.007)	6.04E-07		0.016 (0.014)	2.62E-01		0.032 (0.006)	7.30E-07
6995	X-11301 X-00780			-0.034 (0.007)	1.65E-06		-0.023(0.013)	4.01E-02		-0.032 (0.000)	2.65E-06
Fermented	X-11315			0.034 (0.007)	1.00E 00		0.012 (0.014)	5.02E-01	_	0.029 (0.000)	6 90E-08
dairy	X HOIO			0.004 (0.000)	1.7 12 00		0.000 (0.012)	0.022 01		0.020 (0.000)	0.002 00
Fried	3-phenylpropionate	Amino acid	Phenylalanine &	-0.037 (0.008)	2.82E-06		-0.027 (0.014)	4.79E-02		-0.035 (0.007)	3.59E-07
foods	(hydrocinnamate)		tyrosine metabolism								
	X-11372			0.071 (0.009)	2.09E-15		0.045 (0.010)	3.66E-05		0.060 (0.007)	2.93E-19
	X-11469			-0.035 (0.006)	5.23E-08		-0.009 (0.019)	6.36E-01		-0.032 (0.006)	8.19E-08
	X-02269			-0.035 (0.006)	9.23E-08		-0.008 (0.020)	6.80E-01		-0.032 (0.006)	1.52E-07
	X-11315			-0.042 (0.009)	1.24E-06		-0.022 (0.013)	8.69E-02		-0.036 (0.007)	5.26E-07
Sweets and sweet	docosahexaenoate (DHA; 22:6n3)	Lipid	Essential fatty acid	-0.010 (0.002)	7.56E-09		-0.016 (0.002)	3.97E-09		-0.012 (0.001)	1.07E-17

				Discov	very	Discorda	ant	Meta-an	alysis
Food		Super-							
group	Metabolite	pathway	Sub-pathway	Beta (SE)	Р	Beta (SE)	Р	 Beta (SE)	Р
baked products	eicosapentaenoate (EPA; 20:5n3)	Lipid	Essential fatty acid	-0.009 (0.002)	2.20E-08	-0.013 (0.003)	2.14E-06	-0.010 (0.001)	7.29E-14
	pipecolate	Amino acid	Lysine metabolism	-0.010 (0.002)	1.05E-08	-0.011 (0.004)	4.30E-03	-0.010 (0.002)	9.37E-11
	scyllo-inositol	Lipid	Inositol metabolism	-0.012 (0.002)	7.77E-09	-0.011 (0.005)	3.49E-02	-0.012 (0.002)	5.14E-10
	C- glycosyltryptophan*	Amino acid	Tryptophan metabolism	0.008 (0.002)	3.73E-07	0.008 (0.003)	1.09E-02	0.008 (0.001)	9.24E-09
	glycerate	Carbohydrate	Glycolysis, gluconeogenesis, pyruvate metabolism	-0.011 (0.002)	2.89E-07	-0.006 (0.004)	1.10E-01	-0.009 (0.002)	1.13E-07
	1- docosahexaenoylglyc ero-phosphocholine*	Lipid	Lysolipid	-0.008 (0.002)	3.37E-07	-0.005 (0.004)	1.89E-01	-0.008 (0.001)	1.70E-07
	pyroglutamine*	Amino acid	Glutamate metabolism	0.008 (0.002)	3.05E-06	0.007 (0.004)	9.02E-02	0.007 (0.001)	5.91E-07
	piperine	Xenobiotics	Food component, Plant	-0.009 (0.002)	4.26E-06	-0.007 (0.004)	6.64E-02	-0.008 (0.002)	7.19E-07
	X-11315			-0.013 (0.002)	2.16E-13	-0.004 (0.003)	1.61E-01	-0.011 (0.001)	1.99E-12
	X-11437			-0.010 (0.002)	3.42E-08	-0.002 (0.006)	7.89E-01	-0.009 (0.002)	6.75E-08
	X-12696			0.008 (0.002)	1.84E-06	0.003 (0.004)	4.78E-01	0.007 (0.002)	3.48E-06
Butter and	X-13431	Lipid	Carnitine metabolism	0.034 (0.005)	1.61E-12	0.028 (0.009)	2.67E-03	0.033 (0.004)	6.30E-15
cream	nonanoylcarnitine*								
	myristate (14:0)	Lipid	Long chain fatty acid	0.024 (0.005)	1.04E-06	0.015 (0.011)	1.99E-01	0.022 (0.004)	4.99E-07
	caprate (10:0)	Lipid	Medium chain fatty acid	0.020 (0.004)	4.99E-06	0.017 (0.010)	8.37E-02	0.020 (0.004)	9.38E-07
	X-02249			0.028 (0.005)	1.98E-09	0.025 (0.009)	5.53E-03	0.027 (0.004)	2.22E-11
	X-10510			0.020 (0.004)	3.28E-06	0.022 (0.008)	7.81E-03	0.021 (0.004)	6.04E-08
Spreads	X-11478			0.020 (0.003)	1.69E-08	0.012 (0.005)	1.40E-02	0.017 (0.003)	1.22E-09
and	X-11521			0.017 (0.003)	8.82E-08	0.013 (0.005)	5.17E-03	0.015 (0.003)	1.22E-09
dressings	X-11261			0.020 (0.004)	1.16E-07	0.015 (0.006)	1.15E-02	0.019 (0.003)	3.94E-09
Milk	alpha- hydroxyisovalerate	Amino acid	Valine, leucine and isoleucine metabolism	-0.055 (0.012)	3.85E-06	-0.048 (0.026)	6.36E-02	-0.054 (0.011)	5.48E-07
	X-21365 [trimethyl-N- aminovalerate]			0.107 (0.012)	6.14E-18	0.076 (0.025)	3.08E-03	0.101 (0.011)	2.04E-20
	X-12798			0.093 (0.012)	3.21E-14	0.047 (0.026)	6.94E-02	0.084 (0.011)	7.72E-15
	X-11381			0.076 (0.011)	2.58E-11	0.024 (0.026)	3.54E-01	0.068 (0.010)	5.01E-11
	X-11795			-0.060 (0.012)	3.58E-07	-0.005 (0.021)	8.03E-01	-0.048 (0.010)	3.93E-06
Soy and other milks	glycerate	Carbohydrate	Glycolysis, gluconeogenesis, pyruvate metabolism	0.150 (0.031)	1.14E-06	0.238 (0.048)	2.75E-04	0.175 (0.026)	1.24E-11
Soda	X-11469			-0.020 (0.004)	1.85E-06	-0.001 (0.012)	9.58E-01	-0.018 (0.004)	5.32E-06
Теа	quinate	Xenobiotics	Food component, Plant	-0.019 (0.002)	6.87E-18	-0.025 (0.005)	1.08E-06	-0.020 (0.002)	1.95E-24
	N-acetylornithine	Amino acid	Urea cycle; arginine-, proline-, metabolism	0.009 (0.002)	2.55E-06	0.008 (0.004)	5.49E-02	0.009 (0.002)	3.14E-07
	pseudouridine	Nucleotide	Pyrimidine	0.008 (0.002)	4.01E-06	0.003 (0.005)	5.02E-01	0.008 (0.002)	4.53E-06

				Discov	very		Discorda	ant	_	Meta-an	alysis
Food		Super-	.	_ /	_		_ /	_		_ /	_
group	Metabolite	pathway	Sub-pathway	Beta (SE)	P	,,	Beta (SE)	P		Beta (SE)	<i>P</i>
			metabolism, uracil containing								
	X-14473			-0.026 (0.002)	6.70E-34		-0.021 (0.005)	1.29E-04		-0.025 (0.002)	2.82E-40
	X-12734			0.019 (0.002)	1.32E-20		0.002 (0.006)	7.34E-01		0.018 (0.002)	2.90E-20
	X-09789			0.014 (0.002)	3.40E-13		0.006 (0.005)	2.05E-01		0.013 (0.002)	2.01E-13
	X-14374			-0.012 (0.002)	2.59E-10		-0.008 (0.006)	1.88E-01		-0.012 (0.002)	9.06E-11
Coffee	quinate	Xenobiotics	Food component, Plant	0.035 (0.002)	1.08E-43		0.027 (0.005)	4.30E-08		0.033 (0.002)	1.77E-56
	catechol sulfate	Xenobiotics	Benzoate metabolism	0.017 (0.002)	1.43E-12		0.013 (0.005)	1.84E-02		0.016 (0.002)	4.57E-14
	paraxanthine	Xenobiotics	Xanthine metabolism	0.013 (0.003)	8.63E-07		0.008 (0.005)	1.11E-01		0.012 (0.002)	2.77E-07
	theophylline	Xenobiotics	Xanthine metabolism	0.012 (0.003)	3.88E-06		0.007 (0.005)	1.09E-01		0.011 (0.002)	1.26E-06
	X-14473			0.040 (0.002)	1.53E-77		0.028 (0.005)	6.27E-08		0.039 (0.002)	6.41E-102
	X-14374			0.024 (0.002)	1.98E-28		0.021 (0.006)	1.84E-04		0.023 (0.002)	1.08E-33
	X-05426			0.018 (0.002)	4.64E-13		0.008 (0.005)	9.86E-02		0.016 (0.002)	2.01E-13
	X-12217			0.014 (0.003)	1.64E-07		0.010 (0.006)	1.03E-01		0.013 (0.002)	3.82E-08
	X-09789			-0.014 (0.002)	6.43E-08		-0.004 (0.005)	4.04E-01		-0.012 (0.002)	1.43E-07
	X-12734			-0.014 (0.003)	1.20E-06		-0.007 (0.006)	2.41E-01		-0.013 (0.003)	8.02E-07
Alcohol	scyllo-inositol	Lipid	Inositol metabolism	0.042 (0.005)	2.16E-18		0.037 (0.011)	1.83E-03		0.041 (0.004)	1.57E-21
	alpha- hydroxyisovalerate	Amino acid	Valine, leucine and isoleucine metabolism	0.035 (0.005)	1.29E-12		0.034 (0.009)	4.11E-04		0.035 (0.004)	3.98E-16
	4-androsten- 3beta,17beta-diol disulfate 1*	Lipid	Sterol, Steroid	0.032 (0.004)	1.25E-12		0.037 (0.011)	1.64E-03		0.032 (0.004)	1.89E-15
	eicosapentaenoate (EPA; 20:5n3)	Lipid	Essential fatty acid	0.019 (0.004)	2.26E-07		0.024 (0.005)	7.77E-05		0.021 (0.003)	1.57E-11
	docosahexaenoate (DHA; 22:6n3)	Lipid	Essential fatty acid	0.020 (0.004)	4.03E-08		0.024 (0.007)	1.57E-03		0.020 (0.003)	1.09E-10
	2-oleoylglycero- phosphocholine*	Lipid	Lysolipid	0.016 (0.003)	2.32E-07		0.030 (0.008)	6.93E-04		0.018 (0.003)	6.90E-10
	docosapentaenoate (n3 DPA; 22:5n3)	Lipid	Essential fatty acid	0.019 (0.004)	2.58E-07		0.025 (0.009)	1.15E-02		0.020 (0.003)	7.10E-09
	pipecolate	Amino acid	Lysine metabolism	0.020 (0.004)	6.11E-06		0.021 (0.008)	1.31E-02		0.020 (0.004)	1.71E-07
	1- docosahexaenoylglyc ero-phosphocholine*	Lipid	Lysolipid	0.015 (0.003)	1.39E-06		0.012 (0.009)	1.65E-01		0.015 (0.003)	4.40E-07
	X-11795			0.030 (0.004)	1.11E-11		0.036 (0.009)	1.36E-04		0.031 (0.004)	1.15E-15
	X-10395			0.026 (0.004)	6.35E-11		0.019 (0.010)	6.55E-02		0.025 (0.004)	7.70E-12
	X-14473			0.020 (0.003)	1.06E-09		0.023 (0.009)	1.06E-02		0.020 (0.003)	1.98E-11
	X-12627			0.019 (0.003)	2.06E-08		0.016 (0.008)	5.24E-02		0.018 (0.003)	2.13E-09
	X-11204			0.017 (0.003)	2.34E-07		0.015 (0.008)	5.40E-02		0.017 (0.003)	2.61E-08
	X-11444			-0.018 (0.003)	9.08E-08		-0.012 (0.008)	1.43E-01		-0.017 (0.003)	3.13E-08
	X-12644			0.014 (0.003)	6.55E-06		0.017 (0.008)	2.86E-02		0.014 (0.003)	4.46E-07
	X-11787			-0.014 (0.003)	2.67E-06		-0.007 (0.009)	4.12E-01		-0.014 (0.003)	2.22E-06

Notes: Table shows results of the linear regression analysis for the discovery population (excluding monozygotic twins discordant for each food group), the MZ discordant twin sample and the fixed effects meta-analysis of both groups. Only significant associations are shown which includes those associations passing the bonferroni cut-off in the discovery and fixed effects analyses ($7.60 \times 10^{-6} = 0.05/[20 \text{ food groups } x 329 \text{ detected metabolites}]$) and passing the 5% level of significance in the discordant twin group. The Metabolon platform is a non-targeted platform which identified 456 metabolites in blood for which data were available for 1780 twins.

Superpathway	Metabolite	Food group	beta (SE) P	
Amino acid	3-phenylpropionate (hydrocinnamate)	Fried foods	-0.028 (0.007) 2.44E-05	
	ala ba ha da su da su al sus ta	Whole grains	0.012 (0.003) 6.98E-05	
	aipna-nydroxyisovaierate			
	ninecolate	Willk Sweets and sweet baked products	-0.038 (0.011) 3.60E-04	
	pipecolale	Alcohol	0.016(0.002) 1.76E-00	
	pyroglutamine*	Seafood	-0.064 (0.012) 2.31E-07*	
		Sweets and sweet baked products	0.006 (0.002) 2.35E-04	
Carbohydrate	glycerate	Fruit	0.013 (0.002) 2.38E-11*	
		Soy and other milk	0.132 (0.030) 9.40E-06	
		Sweets and sweet baked products	-0.006 (0.002) 1.97E-04	
Lipid	1-docosahexaenoylglycerophosphocholine*	Seafood	0.083 (0.012) 3.08E-12*	
		AICONOI Sweata and sweat beload products	0.010 (0.003) 1.09E-03	
	docosabezaenoate (DHA: 22:6n3)	Sweets and sweet baked products	-0.004 (0.002) 7.12E-03	
	uocosaliexaelioale (DIIA, 22.015)	Alcohol	0.015 (0.003) 7.79E-08*	
		Sweets and sweet baked products	-0.005 (0.002) 1.71E-03	
		Fruit	0.005 (0.002) 8.25E-03	
	docosapentaenoate (n3 DPA; 22:5n3)	Seafood	0.070 (0.011) 2.94E-10*	
		Alcohol	0.017 (0.003) 1.39E-07*	
	eicosapentaenoate (EPA; 20:5n3)	Seafood	0.112 (0.012) 2.24E-19*	
		Alcohol	0.015 (0.003) 7.21E-08*	
		Whole grains	0.010 (0.003) 7.59E-05	
	scyllo-inositol	Alcohol	-0.004 (0.001) 1.96E-03	
	scyno-mositor	Fruit	0.042 (0.004) 5.05E-22	
		Sweets and sweet baked products	-0.004 (0.002) 3.16E-02	
Nucleotide	pseudouridine	Seafood	-0.065 (0.011) 1.76E-09*	
		Теа	0.007 (0.002) 1.85E-05	
Xenobiotics	quinate	Coffee	0.029 (0.002) 4.90E-35*	
		Теа	-0.012 (0.002) 1.66E-09*	
Unknown	X-02269	Seafood	0.127 (0.012) 6.27E-24*	
		Whole grains	0.012 (0.003) 3.52E-06*	
	Y-00780	Whole grains	-0.014(0.007) $3.75E-02$	
	X-03/03	Tea	0.017(0.003) $3.93E-11$	
		Red, processed meat and eggs	-0.022 (0.006) 3.20E-04	
		Coffee	-0.007 (0.002) 2.48E-03	
	X-11315	Fruit	0.014 (0.002) 1.21E-10*	
		Fermented dairy	0.017 (0.005) 1.17E-03	
		Seafood	0.037 (0.012) 2.03E-03	
		Sweets and sweet baked products	-0.005 (0.002) 2.61E-03	
		vvnole grains	0.007 (0.003) 1.23E-02	
		INUIS and legumes	0.012 (0.006) 3.39E-02	
		Venetable	2.10E-02 2.60E-01	
	X-11372	Fried foods	0.052 (0.007) 9.20E-15*	

Table 2. Multivariate regression results for metabolites associated with multiple food groups

Superpathway	Metabolite	Food group	beta (SE)	Р
		Whole grains	-0.012 (0.003)	1.03E-05
		Fruit	-0.004 (0.002)	4.39E-02
		Vegetable	-0.003 (0.001)	4.85E-02
	X-11381	Milk	0.070 (0.010)	9.83E-12*
		Red, processed meat and eggs	0.035 (0.006)	1.39E-08*
	X-11437	Seafood	0.047 (0.013)	5.05E-04
		Sweets and sweet baked products	-0.006 (0.002)	6.14E-04
	X-11469	Seafood	0.127 (0.012)	2.13E-24*
		Whole grains	0.012 (0.003)	2.89E-06*
		Soda	-0.012 (0.004)	1.74E-03
		Fried foods	-0.014 (0.007)	4.25E-02
	X-11795	Alcohol	0.029 (0.004)	1.48E-13*
		Milk	-0.035 (0.010)	6.93E-04
	X-12627	Alcohol	0.017 (0.003)	9.29E-09*
		Seafood	0.057 (0.011)	1.70E-07*
	X-12644	Seafood	0.067 (0.012)	2.25E-08*
		Alcohol	0.012 (0.003)	3.48E-05
	X-12734	Теа	0.016 (0.002)	2.55E-14*
		Coffee	-0.006 (0.003)	2.38E-02
	X-12798	Milk	0.080 (0.011)	8.80E-13*
		Seafood	-0.042 (0.013)	1.43E-03
	X-14374	Coffee	0.020 (0.002)	2.54E-22*
		Теа	-0.007 (0.002)	3.88E-04
	X-14473	Coffee	0.032 (0.002)	4.05E-57*
		Теа	-0.016 (0.002)	1.47E-17*
		Alcohol	0.011 (0.003)	1.41E-04

Table 2. Multivariate regression	results for metabolites associate	d with multiple food aroups

Notes: Table shows results of a multivariate backwards stepwise linear regression including all food groups associated to a metabolite from the discovery analysis using P<0.05 as the cut-off threshold. Metabolites and foods in bold, and with an asterisk (*) beside the P-value passed the Bonferroni cut-off for multiple testing from the discovery analysis (7.60x10⁻⁶) and were considered significant. The analysis was conducted in 1780 twins.

		Model 1					Standardized	
Food group	Metabolite	R^2	Beta(SE)	Р	R^2	Beta(SE)	Р	betas
Vegetables	X-12063	0.0094	-1.798 (0.627)	4.23E-03				-0.101
Fruit	stachydrine	0.1195	2.327 (0.345)	2.59E-11				0.194
	X-11315		2.586 (0.391)	6.16E-11				0.207
	proline		-1.506 (0.346)	1.48E-05				-0.119
	threitol		1.614 (0.409)	8.63E-05				0.097
	hippurate			5.46E-02				
	glycerate			6.20E-02				
	scyllo-inositol			0.454				
	threonate			0.798				
Whole grains	X-09789	0.0508	1.552 (0.229)	2.08E-11				0.173
	X-11469		1.337 (0.235)	1.79E-08				0.142
	X-02269			0.650				0.173
Nuts and legumes	tryptophan betaine	0.0384	0.905 (0.132)	1.32E-11				0.181
Seafood	docosahexaenoate (DHA; 22:6n3)	0.1514	0.563 (0.090)	7.03E-10	0.1535	0.399 (0.056)	2.09E-12	0.192
	X-11469		0.246 (0.063)	9.44E-05		0.281 (0.061)	3.89E-06	0.130
	pseudouridine		-0.165 (0.047)	5.13E-04		-0.164 (0.046)	4.31E-04	-0.079
	1-eicosatrienoylglycerophosphocholine*		-0.182 (0.067)	6.71E-03		-0.205 (0.063)	1.29E-03	-0.105
	pyroglutamine*		-0.133 (0.050)	8.34E-03		-0.130 (0.049)	8.44E-03	-0.065
	docosapentaenoate (n3 DPA; 22:5n3)		-0.194 (0.076)	1.08E-02				
	1-		-0.110 (0.053)	3.99E-02		-0.123 (0.051)	1.52E-02	
	arachidonoylglycerophosphoethanolami							
	ne*							-0.062
	1-		0.162 (0.081)	4.52E-02		0.169 (0.078)	3.01E-02	
	docosahexaenoylglycerophosphocholine							0.004
				6 275 02				0.084
	elcosapentaenoate (EPA; 20:5n3)			6.37E-02				
	1- linoleovlølvceronhosnhoethanolamine*			7.63E-02				
	1-oleoylglycerophosphoethanolamine			8.13E-02				
	X-02269			0.255				
	3-carboxy-4-methyl-5-propyl-2-			0.271				
	furanpropanoate (CMPF)							
	X-12627			0.450				
	X-12644			0.580				
White meat	3-methylhistidine	0.0291	0.219 (0.034)	1.62E-10				0.170
Red meat	trans-4-hydroxyproline	0.0403	0.449 (0.101)	9.20E-06				0.113
	X-11381		0.393 (0.093)	2.83E-05				0.102
	creatine		0.401 (0.099)	5.40E-05				0.102
Fried foods	X-11372	0.0668	1.058 (0.100)	9.39E-25				0.279
Sweets and sweet baked	C-glycosyltryptophan*	0.0588	2.233 (0.500)	8.93E-06				0.129

Table 3. Backward stepwise linear regression results for foods associated with multiple metabolites and foods associated with single metabolites

		Model 1			Model 2			Standardized
Food group	Metabolite	R ²	Beta(SE)	Р	R ²	Beta(SE)	Р	betas
products	pipecolate		-1.719 (0.439)	9.77E-05				-0 117
	X-12696		1 582 (0 435)	2 94F-04				0.102
	niperine		-1.406 (0.405)	5.40E-04				-0.093
Butter and cream	nonanovlcarnitine*	0.0754	1.030 (0.188)	5.95E-08				0.164
	X-02249		0.615 (0.185)	8.99E-04				0.093
	myristate (14:0)		0.511 (0.175)	3.52E-03				0.083
	caprate (10:0)		0.363 (0.168)	3.05E-02				0.058
	X-10510		0.349 (0.161)	3.09E-02				0.053
Spreads and dressings	X-11261	0.0331	1.103 (0.355)	1.94E-03				0.101
	X-11478		0.774 (0.291)	8.00E-03				0.093
Milk	X-21365 [trimethyl-N-aminovalerate]	0.0735	0.401 (0.065)	1.31E-09				0.175
	X-11381		0.206 (0.059)	4.88E-04				0.091
	X-12798		0.238 (0.069)	6.52E-04				0.102
Теа	X-14473	0.2113	-3.959 (0.503)	1.33E-14				-0.282
	N-acetylornithine		1.710 (0.376)	6.26E-06				0.122
	X-12734		1.675 (0.390)	2.01E-05				0.121
	X-09789		1.461 (0.430)	7.15E-04				0.102
	quinate		-1.253 (0.495)	1.16E-02				-0.090
Coffee	X-14473	0.1963	3.756 (0.404)	2.02E-19				0.344
	quinate		2.197 (0.439)	7.27E-07				0.204
	X-14374			5.46E-02				
	X-05426			0.115				
	catechol sulfate			0.313				
	X-12217			0.715				
	theophylline			0.887				
	paraxanthine			0.964				
Alcohol	scyllo-inositol	0.2537	1.813 (0.213)	1.04E-16	0.2493	1.776 (0.209)	1.16E-16	0.231
	4-androsten-3beta,17beta-diol disulfate		1.692 (0.272)	8.18E-10		1.623 (0.271)	3.21E-09	0.188
	Y_11705		1 107 (0 209)	1 57E-07		1 1/1 (0 206)	1 39F-08	0 1/13
	X-11755 X-11444		-1 130 (0 224)	5.74E-07		-1 075 (0 221)	1.33E-06	-0 131
	alpha-hydroxyisovalerate		1.066 (0.224)	2 28E-06		1.010 (0.221)	2 54E-06	0.131
	eicosapentaenoate (EPA: 20:5n3)		1.000 (0.221)	2.23E-04		0.827 (0.222)	2.06E-04	0.098
	2-oleoylglycerophosphocholine*		0.630 (0.206)	2.30E-03		0.626 (0.203)	2.11E-03	0.080
	X-11787		-0.491 (0.217)	2.42E-02		-0.442 (0.217)	4.18E-02	-0.053
	X-12627		-0.484 (0.241)	4.51E-02				
	X-10395			0.112				
	docosapentaenoate (n3 DPA: 22:5n3)			0.489				
	docosahexaenoate (DHA; 22:6n3)			0.502				
	X-11204			0.928				

Table 3. Backward stepwise linear regression results for foods associated with multiple metabolites and foods associated with single metabolites

Notes: Table shows the results of a multivariate backwards stepwise linear regression including all associated metabolites used to predict food group intakes using *P*<0.05 as the cut-off threshold. Metabolites highlighted in red indicate the association was in the opposite direction to the discovery analysis for model 1 and therefore, for the corresponding food group a second multivariate regression was undertaken. The rightmost column shows the final standardized betas used for the weighted scoring method. The analysis was conducted in 1780 twins.

			Training a	group	Test gro	up
Food group	No. metabolites in score	Score	Beta (SE)	Р	Beta (SE)	Р
Vegetables	1	Top: X-12063	-0.005 (0.001)	2.96E-06	-0.006 (0.002)	3.62E-04*
		Quartiles	0.004 (0.001)	8.59E-03	0.005 (0.002)	6.88E-03*
Fruit	4	Top: X-11315	0.019 (0.002)	7.14E-20	0.016 (0.002)	1.81E-18
		Quartiles	0.064 (0.005)	4.06E-34	0.062 (0.005)	1.28E-36
		Continuous	0.060 (0.005)	2.68E-36	0.055 (0.004)	3.76E-37
		Weighted	0.010 (0.001)	3.03E-36	0.009 (0.001)	4.48E-40
Whole grains	3	Top: X-09789	0.020 (0.003)	1.33E-13	0.031 (0.003)	4.35E-22
		Quartiles	0.038 (0.005)	6.99E-16	0.049 (0.005)	5.42E-23
		Continuous	0.036 (0.004)	1.40E-18	0.046 (0.005)	6.98E-22
		Weighted	0.006 (0.001)	8.78E-19	0.008 (0.001)	9.51E-23
Nuts and legumes	1	Top: tryptophan betaine	0.044 (0.006)	1.45E-13	0.037 (0.005)	3.38E-13
		Quartiles	0.041 (0.005)	2.78E-14	0.037 (0.005)	1.29E-13
Seafood	7	Top: DHA	0.147 (0.013)	6.98E-30	0.141 (0.011)	1.76E-34
		Quartiles	0.711 (0.048)	1.24E-45	0.599 (0.049)	7.85E-32
		Continuous	0.662 (0.046)	1.30E-42	0.575 (0.045)	3.43E-34
		Weighted	0.076 (0.005)	6.01E-44	0.069 (0.005)	1.63E-41
White meat	1	Top: 3-methylhistidine	0.135 (0.021)	1.52E-10	0.077 (0.023)	7.78E-04
		Quartiles	0.176 (0.021)	2.11E-16	0.134 (0.022)	1.03E-09
Red meat	3	Top: trans-4-hydroxyproline	0.035 (0.006)	9.27E-08	0.036 (0.007)	3.66E-08
		Quartiles	0.108 (0.013)	9.64E-16	0.124 (0.014)	9.23E-19
		Continuous	0.100 (0.012)	2.96E-16	0.114 (0.012)	3.43E-19
		Weighted	0.011 (0.001)	3.40E-16	0.012 (0.001)	3.79E-19
Fried foods	1	Top: X-11372	0.067 (0.007)	3.28E-20	0.080 (0.009)	2.55E-17
		Quartiles	0.068 (0.008)	2.30E-18	0.084 (0.010)	1.31E-16
Sweets and sweet baked products	4	Top: C-glycosyltryptophan*	0.136 (0.034)	8.54E-05	0.008 (0.001)	1.78E-09
		Quartiles	0.457 (0.095)	2.00E-06	0.032 (0.004)	6.44E-17
		Continuous	0.427 (0.082)	2.27E-07	0.031 (0.003)	8.93E-21
		Weighted	0.047 (0.009)	1.73E-07	0.003 (0.000)	3.28E-21
Butter and creams	5	Top: nonanoylcarnitine*	0.034 (0.004)	2.69E-16	0.022 (0.004)	8.92E-08
		Quartiles	0.122 (0.013)	2.53E-21	0.107 (0.013)	4.89E-16
		Continuous	0.120 (0.013)	1.90E-18	0.094 (0.012)	8.66E-15
		Weighted	0.011 (0.001)	8.09E-21	0.009 (0.001)	2.30E-15
Spreads and dressings	2	Top: X-11261	0.019 (0.003)	2.77E-09	0.014 (0.003)	6.34E-06
		Quartiles	0.041 (0.006)	4.21E-12	0.034 (0.007)	4.86E-07
		Continuous	0.036 (0.005)	5.47E-11	0.026 (0.006)	1.83E-05*
		Weighted	0.003 (0.001)	5.89E-11	0.002 (0.001)	1.68E-05*
Milk	3	Top: X-21365	0.102 (0.011)	2.58E-20	0.094 (0.010)	6.20E-19
		Quartiles	0.264 (0.026)	3.02E-23	0.247 (0.028)	1.60E-18
		Continuous	0.250 (0.023)	1.23E-25	0.204 (0.024)	4.95E-17
		Weighted	0.032 (0.003)	2.90E-26	0.027 (0.003)	1.49E-19
Теа	5	Top: X-14473	-0.025 (0.002)	2.56E-37	-0.024 (0.002)	2.83E-36
		Quartiles	0.078 (0.005)	3.98E-42	0.050 (0.005)	3.32E-22
		Continuous	0.073 (0.005)	2.96E-50	0.051 (0.004)	2.08E-32
		Weighted	0.012 (0.001)	1.15E-52	0.009 (0.001)	7.64E-38
		-	. ,		. ,	

Table 4. Associations between reported food intakes and the final scores in the training and test groups

			Training g	roup	Test grou	лр
Food group	No. metabolites in score	Score	Beta (SE)	Р	Beta (SE)	Р
Coffee	2	Top: X-14473	0.038 (0.002)	3.47E-83	0.036 (0.002)	3.00E-61
		Quartiles	0.090 (0.003)	8.56E-115	0.091 (0.004)	1.06E-104
		Continuous	0.064 (0.003)	2.63E-82	0.057 (0.003)	1.66E-71
		Weighted	0.018 (0.001)	4.83E-87	0.016 (0.001)	5.92E-72
Alcohol	8	Top: scyllo-inositol	0.041 (0.004)	8.22E-22	0.039 (0.004)	2.48E-24
		Quartiles	0.212 (0.018)	7.53E-31	0.191 (0.013)	1.21E-42
		Continuous	0.197 (0.018)	1.06E-27	0.180 (0.013)	2.51E-41
		Weighted	0.029 (0.003)	5.47E-27	0.027 (0.002)	7.79E-45

Table 4. Associations between reported food intakes and the final scores in the training and test groups

Notes: A linear regression was performed in the training (n=1780) and test groups (n=1779) using each food group intake as a predictor of the top metabolite and metabolite scores. An asterisk (*) beside the *P*-value indicates the association does not pass the Bonferroni cut-off from the discovery analysis (P<7.60x10⁻⁶)

	Tertile 1	Tertile 3
Food group	(<i>n</i> =593)	(<i>n</i> =593)
Vegetables	19.5 (5.4)	51.0 (13.4)
Fruit	9.7 (4.4)	35.6 (10.3)
Whole grains	2.7 (2.4)	19.2 (7.4)
Nuts and legumes	3.0 (1.5)	12.9 (5.4)
Seafood	0.7 (0.5)	4.5 (1.8)
White meat	0.6 (0.4)	3.4 (0.8)
Red meat	2.9 (1.5)	10.8 (3.0)
Fried foods	1.6 (1.1)	8.0 (3.3)
Sweets and sweet baked products	4.2 (3.6)	30.4 (14.5)
Butter and cream	-0.4 (1.0)	10.6 (7.4)
Spreads and dressings	1.7 (2.1)	17.3 (10.8)
Milk	1.1 (0.9)	5.8 (1.5)
Теа	3.2 (3.7)	35.5 (5.6)
Coffee	0.0 (0.5)	22.1 (7.9)
Alcohol	0.4 (1.0)	14.3 (8.8)

Table 5. Average (SD) energy-adjusted intakes (servings/week) for the top and bottom tertiles for each food in the test group

Table 6. Results of ROC analysis

						Versus t	op metabolite
Food group	Score	Sensitivity	Specificity	Correctly classified	ROC area	y ²	Р
Vegetables	Top: X-12063	59.74%	55.47%	57.60%	0.6136 [0.5802, 0.6470]		
	Quartiles	59.22%	54.50%	56.85%	0.6136 [0.5802, 0.6470]	0.00	1.0000
Fruit	Top: X-11315	65.59%	60.34%	62.98%	0.6783 [0.6479, 0.7087]		
	Quartiles	68.54%	67.06%	67.80%	0.7305 [0.7020, 0.7590]	18.14	< 0.0001
	Continuous	67.52%	66.55%	67.04%	0.7339 [0.7057. 0.7622]	19.99	< 0.0001
	Weighted	66.84%	67.24%	67.04%	0.7411 [0.7131. 0.7690]	29.77	< 0.0001
Whole grains	Top: X-09789	65.25%	67.07%	66.15%	0.7134 [0.6841. 0.7428]		
	Quartiles	66.61%	65.47%	66.04%	0.7063 [0.6768. 0.7359]	0.60	0.4394
	Continuous	67.46%	66.32%	66.89%	0.7186 [0.6895, 0.7478]	0.36	0.5500
	Weighted	67.63%	65.81%	66.72%	0.7224 [0.6934, 0.7514]	1.37	0.2425
Nuts and legumes	Top: tryptophan betaine	64.25%	60.70%	62.47%	0.6723 [0.6366, 0.7080]		
•	Quartiles	65.82%	57.68%	61.77%	0.6629 [0.6268, 0.6991]	2.42	0.1197
Seafood	Top: DHA	64.79%	67.46%	66.13%	0.7230 [0.6943, 0.7516]		
	Quartiles	65.81%	67.12%	66.47%	0.7307 [0.7023, 0.7592]	0.42	0.5172
	Continuous	66.15%	67.12%	66.64%	0.7356 [0.7075, 0.7637]	1.12	0.2893
	Weighted	69.57%	70.68%	70.13%	0.7547 [0.7274, 0.7821]	11.88	0.0006
White meat	Top: 3-methylhistidine	77.78%	32.44%	56.86%	0.5733 [0.5371, 0.6096]		
	Quartiles	54.31%	62.39%	58.40%	0.5536 [0.5172, 0.5901]	4.74	0.0295
Red meat	Top: trans-4-hydroxyproline	58.94%	65.37%	62.16%	0.6697 [0.6390, 0.7003]		
	Quartiles	61.73%	63.50%	62.62%	0.6920 [0.6622, 0.7218]	5.36	0.0207
	Continuous	63.10%	65.70%	64.40%	0.6963 [0.6666, 0.7260]	8.63	0.0033
	Weighted	63.61%	65.53%	64.57%	0.6963 [0.6666, 0.7260]	9.36	0.0022
Fried foods	Top: X-11372	65.30%	66.32%	65.81%	0.7322 [0.7038, 0.7606]		
	Quartiles	65.98%	65.93%	65.96%	0.7217 [0.6928, 0.7506]	6.70	0.0096
Sweets and sweet baked products	Top: C-glycosyltryptophan*	58.67%	56.51%	57.59%	0.6002 [0.5679, 0.6325]		
	Quartiles	58.57%	57.78%	58.18%	0.6235 [0.5917, 0.6552]	3.29	0.0697
	Continuous	60.27%	59.32%	59.80%	0.6249 [0.5932, 0.6566]	4.29	0.0384
	Weighted	60.10%	58.97%	59.54%	0.6254 [0.5937, 0.6571]	5.07	0.0243
Butter and cream	Top: nonanoylcarnitine*	71.88%	39.55%	56.84%	0.5889 [0.5529, 0.6249]		
	Quartiles	61.02%	58.87%	59.95%	0.6386 [0.6037, 0.6735]	7.45	0.0064
	Continuous	58.31%	60.75%	59.52%	0.6383 [0.6034, 0.6732]	7.38	0.0066
	Weighted	59.15%	59.90%	59.52%	0.6385 [0.6036, 0.6734]	11.39	0.0007
Spreads and dressings	Top: X-11261	59.08%	57.48%	58.28%	0.6199 [0.5880, 0.6518]		
	Quartiles	60.27%	60.10%	60.19%	0.6374 [0.6059, 0.6689]	5.42	0.0199
	Continuous	59.25%	60.61%	59.93%	0.6319 [0.6003, 0.6636]	3.07	0.0799
	Weighted	58.90%	60.27%	59.59%	0.6316 [0.5999, 0.6633]	3.14	0.0762
Milk	Top: X-21365	60.17%	60.65%	60.41%	0.6409 [0.6096, 0.6723]		
	quartiles	60.51%	61.33%	60.92%	0.6403 [0.6089, 0.6717]	0.00	0.9575
	Continuous	60.68%	61.33%	61.00%	0.6355 [0.6040, 0.6669]	0.22	0.6402
T	Weighted	60.68%	60.48%	60.58%	0.6442 [0.6130, 0.6755]	0.14	0.7122
lea	Top: X-14473	59.28%	73.51%	67.04%	0.7384 [0.7078, 0.7691]	0.00	0.1012
	Quartiles	61.63%	64.63%	63.13%	0.7121 [0.6803, 0.7439]	2.68	0.1016
	Continuous	66.04%	66.16%	66.10%	0.7257 [0.6943, 0.7572]	0.80	0.3705
	Weighted	67.40%	65.65%	66.53%	0.7443 [0.7139, 0.7747]	0.40	0.5276

Table 6. Results of ROC analysis

						Versus t	op metabolite
Food group	Score	Sensitivity	Specificity	Correctly classified	ROC area	χ²	Р
Coffee	Top: X-14473	78.21%	65.38%	72.42%	0.7779 [0.7483, 0.8074]		
	Quartiles	72.06%	76.36%	74.21%	0.7917 [0.7627, 0.8208]	3.08	0.0790
	Continuous	66.78%	70.92%	68.85%	0.7803 [0.7510, 0.8096]	0.11	0.7388
	Weighted	67.63%	70.75%	69.19%	0.7856 [0.7565, 0.8148]	2.06	0.1511
Alcohol	Top: scyllo-inositol	75.75%	63.32%	70.02%	0.7531 [0.7230, 0.7832]		
	Quartiles	71.48%	70.97%	71.22%	0.7914 [0.7638, 0.8191]	5.52	0.0188
	Continuous	70.80%	74.02%	72.41%	0.8067 [0.7800, 0.8333]	11.05	0.0009
	Weighted	74.70%	76.40%	75.55%	0.8351 [0.8104, 0.8598]	38.99	<0.0001

Notes: Table shows the results of the receiver operating characteristic analysis testing the ability of each top metabolite or metabolite score to predict the top (1, positive outcome) and bottom (0, negative outcome) tertiles of food group intake. The area under the receiver operating characteristic curve (AUC) was compared for each metabolite score against the top performing metabolite. Metabolite scores which performed the best and better than the top metabolite are in bold. The analysis was conducted in the test group (n=1779).

Table 7. Top metabolite and score associations with food preferences in the test group

			Liking		
	No.				
	metabolites in				
Food group	score	Score	Beta (SE)	Р	
Vegetables	1	Top: X-12063	-0.062 (0.059)	NS	
		Quartiles	0.036 (0.067)	NS	
Fruit	4	Top: X-11315	0.203 (0.045)	7.40E-06	
		Quartiles	0.490 (0.122)	6.62E-05	
		Continuous	0.448 (0.108)	3.51E-05	
		Weighted	0.077 (0.017)	1.12E-05	
Whole grains	3	Top: X-09789	0.287 (0.054)	1.37E-07	
		Quartiles	0.226 (0.083)	6.62E-03*	
		Continuous	0.258 (0.075)	6.29E-04	
		Weighted	0.045 (0.012)	1.70E-04	
Nuts and legumes	1	Top: tryptophan betaine	0.271 (0.045)	3.04E-09	
	_	Quartiles	0.293 (0.042)	9.87E-12	
Seafood	/	Top: DHA	0.333 (0.035)	6.85E-20	
		Quartiles	1.538 (0.139)	2.12E-26	
		Continuous	1.464 (0.129)	1.01E-27	
White most	1	Weighted	0.173 (0.014)	1.51E-30	
white meat	L	Top: 3-methymistidine	0.083 (0.037)	2.45E-02	
Pod most	2	Ton: trans-4-bydroxyproling	0.118 (0.055)	0.20E-04	
Red meat	3	Ouartilos	0.221 (0.040)	2.29E-00	
		Continuous	0.834 (0.093)	4.01E-19 3.5/E-19	
		Weighted	0.081 (0.009)	7 71F-19	
Fried foods	1	Top: X-11372	0.031(0.009)	2 69F-13	
Theu loous	-	Quartiles	0.355 (0.051)	1 12F-11	
Sweets and sweet baked	4	Top: C-glycosyltryptophan*	0.004 (0.002)	1.09E-02*	
products					
		Quartiles	0.021 (0.004)	5.24E-07	
		Continuous	0.020 (0.004)	2.07E-07	
		Weighted	0.002 (0.000)	2.24E-07	
Butter and creams	5	Top: nonanoylcarnitine*	0.023 (0.033)	4.79E-01*	
		Quartiles	0.294 (0.094)	1.78E-03	
		Continuous	0.262 (0.080)	1.11E-03	
<u> </u>		Weighted	0.022 (0.007)	2.61E-03	
Spreads and dressings	2	Top: X-11261	0.105 (0.041)	1.15E-02*	
		Quartiles	0.207 (0.089)	2.06E-02*	
		Weighted	0.203 (0.073)	5.71E-05	
Milk	2	Top: V-21265	0.020 (0.007)	5.07E-05	
	5	Ouartiles	0.384 (0.116)	9.555-04	
		Continuous	0.352 (0.097)	3 15F-04	
		Weighted	0.047 (0.012)	9 71E-05	
Теа	5	Top: X-14473	-0.217 (0.031)	7.83E-12	
	5	Quartiles	0.624 (0.082)	8.88E-14	
		Continuous	0.549 (0.068)	3.19E-15	
		Weighted	0.089 (0.011)	2.53E-16	
Coffee	2	Top: X-14473	0.241 (0.055)	1.36E-05	
		Quartiles	0.689 (0.101)	1.98E-11	
		Continuous	0.379 (0.073)	2.50E-07	
		Weighted	0.104 (0.020)	4.38E-07	
Alcohol	8	Top: scyllo-inositol	0.322 (0.042)	1.01E-13	
		Quartiles	1.559 (0.144)	1.22E-25	
		Continuous	1.299 (0.125)	6.46E-24	
		Weighted	0.197 (0.018)	4.08E-27	

NS, not significant; *, not statistically significant (P<3.33x10⁻³). Notes: Table shows the results of the linear regression using each food group liking score used to predict the top metabolite and metabolite score. The analysis was conducted in the test group (*n*=1779).

Appendix E. Chapter 6 Appendices

		Hippurate		Diet score		Foods (3)
OTU ID ⁽²⁾	Assigned taxonomy	Beta (SE)	Р	Beta (SE)	Р	(<i>P</i> <0.05)
denovo467	k_Bacteria; p_Actinobacteria; c_Actinobacteria; o_Actinomycetales;	-0.099 (0.022)	5.14E-06	-0.051 (0.011)	2.81E-06	Fruit: -0.005(0.002)
denovo44	k_Bacteria; p_Bacteroidetes; c_Bacteroidia; o_Bacteroidales; f_Rikenellaceae;	0.094 (0.023)	6.55E-05			110. 0.000(0.000)
denovo299	y, s k_Bacteria; pFirmicutes; cClostridia; oClostridiales; f; g; s	0.113 (0.024)	2.21E-06	0.044 (0.010)	9.76E-06	Coffee: 0.013(0.002)*
denovo346	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_; g_; s_	0.129 (0.025)	2.07E-07			
denovo197	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Clostridiaceae; g_; s	0.126 (0.026)	1.44E-06			
denovo52	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Clostridiaceae; a_Clostridium: s	0.117 (0.024)	1.50E-06			
denovo1263	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae	-0.097 (0.022)	1.38E-05			
denovo272	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae;	-0.094 (0.022)	1.29E-05			
denovo20	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae;	-0.117 (0.022)	1.29E-07			
denovo100	kBacteria; pFirmicutes; cClostridia; oClostridiales; fLachnospiraceae;	-0.107 (0.022)	1.92E-06			
denovo55	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; a_Riminococcus]: s	-0.123 (0.021)	1.17E-08	-0.054 (0.011)	2.79E-06	Fruit: -0.006(0.002)*
denovo27	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; d_IRuminococcus); s_maxus	-0.107 (0.023)	3.04E-06	-0.064 (0.011)	1.99E-08	Fruit: -0.006(0.002)*
denovo25	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; Bautia: s	-0.135 (0.024)	1.28E-08			
denovo35	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; 	-0.137 (0.022)	4.94E-10			
denovo13	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; g_Blautia: s	-0.177 (0.021)	5.56E-17			
denovo144	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; g_Blautia: s	-0.165 (0.022)	4.83E-14			
denovo237	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; g_Blautia; s_producta	-0.102 (0.022)	4.78E-06			
denovo329	K_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; g_Dorea: s	-0.130 (0.023)	1.73E-08			
denovo97	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; g_Dorea: s_formicigenerans	-0.114 (0.022)	2.46E-07			
denovo414	K_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Lachnospiraceae; g_Roseburia; s	-0.091 (0.022)	3.67E-05			
denovo79	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae;	0.097 (0.022)	8.32E-06			
denovo123	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_:s	0.103 (0.024)	2.83E-05			
denovo12	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_:s	0.167 (0.025)	8.61E-11			
denovo1380	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae;	0.095 (0.022)	1.10E-05			

Table 1. List of OTUs associated with hippurate, the hippurate diet score and foods ⁽¹⁾

		Hippura	ate	Diet sc	ore	Foods ⁽³⁾
OTU ID ⁽²⁾	Assigned taxonomy	Beta (SE)	Ρ	Beta (SE)	Ρ	(<i>P</i> <0.05)
denovo438	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_; s_	0.100 (0.025)	5.01E-05			
denovo33	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_; s_	0.098 (0.024)	4.87E-05			
denovo469	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_Faecalibacterium; s_prausnitzii	0.100 (0.023)	1.66E-05	0.034 (0.010)	9.24E-04	WG: 0.007(0.003)
denovo276	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_Oscillospira; s_	0.110 (0.026)	2.53E-05			
denovo59	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_Oscillospira; s_	0.170 (0.026)	2.17E-10			
denovo424	k_Bacteria; p_Firmicutes; c_Clostridia; o_Clostridiales; f_Ruminococcaceae; g_Ruminococcus: s	0.107 (0.026)	3.36E-05			

Table 1. List of OTUs associated with hippurate, the hippurate diet score and foods ⁽¹⁾

*= statistically significant: P<0.0017; WG: whole grain products; OTUs, operational taxonomic units

(4) Microbiome OTUs significantly associated with both hippurate and the hippurate diet score are shown. Associations were adjusted for covariates (age, Shannon Index, metabolite batch, BMI, sex and family relatedness) and multiple testing using Bonferroni correction. Hippurate diet score associations were also adjusted for hippurate.

(5) OTU ID assignment is specific to the TwinsUK cohort.

(6) All foods included in the hippurate diet score were fitted into a backwards stepwise linear regression using P<0.05 as the cut-off threshold with each collapsed taxa/OTU associated to both hippurate and the diet score. Results displayed are the betas with standard errors of foods at least nominally associated (P<0.05). Statistical significance was defined as P<0.0017 (Bonferronil: 0.05/[10 taxa x 3 foods]).</p>

					Hippura	te	Diet score		Foods
Phylum	Class	Order	Family	Genus	Beta (SE)	Р	Beta (SE)	Р	P<0.05 ⁽²⁾
Actinobacteria	Actinobacteria	Actinomycetales			-0.083 (0.022)	1.31E-04	-0.035 (0.011)	1.67E-03	Fruit: -0.004(0.002) WG: -0.007(0.003)
Actinobacteria	Actinobacteria	Actinomycetales	Actinomycetaceae		-0.089 (0.021)	2.89E-05	-0.036 (0.011)	1.70E-03	Fruit: -0.004(0.002) WG: -0.007(0.003)
Actinobacteria	Actinobacteria	Actinomycetales	Actinomycetaceae	Actinomyces	-0.101 (0.021)	1.55E-06	-0.045 (0.011)	5.71E-05	Fruit: -0.005(0.002) WG: -0.008(0.003)
Firmicutes					-0.102 (0.024)	1.94E-05			. ,
Firmicutes	Clostridia				-0.088 (0.024)	2.44E-04			
Firmicutes	Clostridia	Clostridiales			-0.088 (0.024)	2.40E-04			
Firmicutes	Clostridia	Clostridiales	Lachnospiraceae		-0.125 (0.022)	1.72E-08			
Firmicutes	Clostridia	Clostridiales	Lachnospiraceae	[Ruminococcus]	-0.111 (0.022)	4.03E-07	-0.038 (0.011)	6.35E-04	Fruit: -0.005(0.002) WG: -0.008(0.003)
Firmicutes	Clostridia	Clostridiales	Lachnospiraceae	Blautia	-0.146 (0.021)	5.68E-12			· · · ·
Firmicutes	Clostridia	Clostridiales	Lachnospiraceae	Dorea	-0.116 (0.021)	6.13E-08			
Firmicutes	Clostridia	Clostridiales	Lachnospiraceae	Roseburia	-0.089 (0.022)	4.18E-05			
Firmicutes	Erysipelotrichi				-0.111 (0.021)	1.64E-07			
Firmicutes	Erysipelotrichi	Erysipelotrichales			-0.111 (0.021)	1.64E-07			
Firmicutes	Erysipelotrichi	Erysipelotrichales	Erysipelotrichaceae		-0.111 (0.021)	1.64E-07			
Firmicutes	Erysipelotrichi	Erysipelotrichales	Erysipelotrichaceae	[Eubacterium]	-0.083 (0.021)	9.30E-05	-0.040 (0.012)	6.12E-04	Fruit: -0.004(0.002) WG: -0.010(0.003)*
Proteobacteria	Betaproteobacteria	Burkholderiales	Oxalobacteraceae	Ralstonia	-0.084 (0.021)	7.85E-05			. ,

Table 2. List of collapsed taxa associated with hippurate, the hippurate diet score and foods ⁽¹⁾

*= statistically significant: *P*<0.0017; WG: whole grain products

(1) Microbiome collapsed taxa significantly associated with both hippurate and the hippurate diet score are shown. Associations were adjusted for covariates (age, Shannon Index, metabolite batch, BMI, sex and family relatedness) and multiple testing using Bonferroni correction. Hippurate diet score associations were also adjusted for hippurate.

(2) All foods included in the hippurate diet score were fitted into a backwards stepwise linear regression using *P*<0.05 as the cut-off threshold with each collapsed taxa associated to both hippurate and the diet score. Results displayed are the betas with standard errors of foods at least nominally associated (*P*<0.05). Statistical significance was defined as *P*<0.0017 (Bonferronil: 0.05/[10 taxa/OTUs x 3 foods])

Table 3. Associations between MetS status and criteria categories with Shannon diversity, the hippurate gradient, the hippurate diet score and associated OTUs and taxa

MetS				
phenotype	Variable	Beta(SE)	Р	R ²
MetS status	Shannon diversity	OR: 0.744 (0.064)	6.35E-04	0.0105
(0, no; 1, yes)	Hippurate trajectory	OR: 0.795 (0.082)	0.026	0.0054
	Actinomycetaceae family	OR: 1.397 (0.145)	0.001	0.0110
	Actinomycetales order	OR: 1.373 (0.140)	0.002	0.0100
	Actinomyces genus	OR: 1.310 (0.134)	0.009	0.0072
	Eubacterium genus	OR: 1.215 (0.111)	0.034	0.0038
	Ruminococcus genus	OR: 1.308 (0.128)	0.006	0.0075
	Ruminococcus genus OTU	OR: 1.373 (0.145)	0.003	0.0099
	denovo55			
	Clostridiales order OTU denovo299	OR: 0.703 (0.063)	9.47E-05	0.0122
	Faecalibacterium prausnitzii OTU denovo469	OR: 0.732 (0.065)	4.33E-04	0.0095
BMI	Shannon diversity	-0.770 (0.151)	4.63E-07	0.0288
	Hippurate trajectory	-0.700 (0.149)	3.25E-06	0.0240
	Hippurate diet	-0.332 (0.154)	0.032	0.0053
	Actinomycetaceae family	0.393 (0.152)	0.010	0.0075
	Actinomycetales order	0.420 (0.153)	0.006	0.0085
	Actinomyces genus	0.431 (0.155)	0.005	0.0089
	Ruminococcus genus	0.407 (0.149)	0.006	0.0081
	Ruminococcus genus OTU denovo55	0.430 (0.151)	0.005	0.0091
	Clostridiales order OTU denovo299	-0.317 (0.147)	0.031	0.0048
	Faecalibacterium prausnitzii OTU denovo469	-0.487 (0.143)	7.26E-04	0.0115
HDL-	Shannon diversity	0.096 (0.027)	4.42E-04	0.0126
cholesterol	Hippurate diet	0.081 (0.027)	0.003	0.0095
	Ruminococcus genus OTU denovo55	0.092 (0.027)	8.46E-04	0.0123
	Clostridiales order OTU denovo299	0.073 (0.027)	0.007	0.0076
	Faecalibacterium prausnitzii OTU denovo469	0.056 (0.026)	0.035	0.0044
Triglycerides	Shannon	-0.122 (0.027)	4.97E-06	0.0213
	Hippurate	-0.077 (0.026)	0.003	0.0087
	Diet	-0.094 (0.027)	5.72E-04	0.0130
	Actinomycetaceae family	0.055 (0.025)	0.029	0.0045
	Eubacterium genus	0.061 (0.026)	0.019	0.0056
	Ruminococcus genus	0.072 (0.026)	0.005	0.0077
	Ruminococcus genus OTU	0.109 (0.026)	3.56E-05	0.0178
	denovo55			
	Clostridiales order OTU denovo299	-0.115 (0.024)	2.38E-06	0.0194
	Faecalibacterium prausnitzii OTU denovo469	-0.079 (0.025)	0.002	0.0092

MetS, metabolic syndrome; HDL, high density lipoprotein; OTU, operational taxonomic unit; OR, odds ratio Notes: A linear regression was performed using Shannon diversity, the hippurate trajectory, the hippurate diet and hippurate and diet associated taxa/OTUs as predictors of MetS status (adjusting for age, sex, and family relatedness) and each component adjusting for age, BMI (except for BMI), sex, and family relatedness. Statistical significance was defined as *P*<0.05.

				%
		Hipp	variance	
MetS		traje	ctory	through
phenotype	Variable ⁽¹⁾	r_{x}^{2} ⁽²⁾	r_{xy}^{2} (3)	variable
MetS status	Hippurate trajectory	0.0054		
(0, no; 1, yes)	Shannon diversity		0.0021	61.1
	Actinomycetaceae family		0.0040	25.9
	Actinomycetales order		0.0042	22.2
	Actinomyces genus		0.0042	22.2
	Eubacterium genus		0.0052	3.7
	Ruminococcus genus		0.0046	14.8
	Ruminococcus genus OTU denovo55		0.0038	29.6
	Clostridiales order OTU denovo299		0.0026	51.9
	Faecalibacterium prausnitzii OTU denovo469		0.0031	42.6
BMI	Hippurate trajectory	0.0288		
	Hippurate diet		0.0203	29.5
	Shannon diversity		0.0122	57.6
	Actinomycetaceae family		0.0215	25.3
	Actinomycetales order		0.0217	24.7
	Actinomyces genus		0.0211	26.7
	Ruminococcus genus		0.0223	22.6
	Ruminococcus genus OTU denovo55		0.0207	28.1
	Clostridiales order OTU denovo299		0.0199	30.9
	Faecalibacterium prausnitzii OTU denovo469		0.0186	35.4
Triglycerides	Hippurate trajectory	0.0087		
	Hippurate diet		0.0055	36.8
	Shannon diversity		0.0032	63.2
	Actinomycetaceae family		0.0077	11.5
	Eubacterium genus		0.0084	3.4
	Ruminococcus genus		0.0079	9.2
	Ruminococcus genus OTU denovo55		0.0063	27.6
	Clostridiales order OUT denovo299		0.0045	48.3
	Faecalibacterium prausnitzii OTU denovo469		0.0060	31.0

 Table 4. Percent variance in the association between the hippurate trajectory and MetS,

 BMI and triglycerides accounted for through each associated variable

MetS, metabolic syndrome; OTU, operational taxonomic unit

(5) All variables were significantly (P<0.05) associated with each listed MetS phenotype.

(6) The proportion of the variance in each MetS phenotype explained by the hippurate trajectory after taking into account all covariates (age, sex, family relatedness and BMI [for triglycerides]).

(7) The proportion of the variance in each MetS phenotype explained by the hippurate trajectory after taking into account all covariates as in (2) and adjusting for the applicable variable.

		Hippurate diet		% variance	
		SC	ore	through	
MetS phenotype	Variable ⁽¹⁾	$r_{x}^{(2)}$	r ² _{xy} (3)	variable	
BMI	Hippurate diet	0.0053			
	Shannon diversity		0.0029	45.3	
	Hippurate trajectory		0.0021	60.4	
	Actinomycetaceae family		0.0042	20.8	
	Actinomycetales order		0.0042	20.8	
	Actinomyces genus		0.0039	26.4	
	Ruminococcus genus		0.0045	15.1	
	Ruminococcus genus OTU denovo55		0.0038	28.3	
	Clostridiales order OTU denovo299		0.0038	28.3	
	Faecalibacterium prausnitzii OTU denovo469		0.0034	35.8	
HDL	Hippurate diet	0.0095			
	Shannon diversity		0.0074	22.1	
	Ruminococcus genus OTU denovo55		0.0073	23.2	
	Clostridiales order OTU denovo299		0.0070	26.3	
	Faecalibacterium prausnitzii OTU denovo469		0.0079	16.8	
Triglycerides	Hippurate diet	0.0130			
	Shannon diversity		0.0100	23.1	
	Hippurate trajectory		0.0097	25.4	
	Actinomycetaceae family		0.0118	9.2	
	Eubacterium genus		0.0118	9.2	
	Ruminococcus genus		0.0120	7.7	
	Ruminococcus genus OTU denovo55		0.0100	23.1	
	Clostridiales order OUT denovo299		0.0087	33.1	
	Faecalibacterium prausnitzii OTU denovo469		0.0104	20.0	

 Table 5. Percent variance in the association between the hippurate diet score and BMI,

 HDL cholesterol and triglycerides accounted for through each associated variable

MetS, metabolic syndrome; HDL, high density lipoprotein; OTU, operational taxonomic unit

(1) All variables were significantly (P<0.05) associated with each listed MetS phenotype.

(2) The proportion of the variance in each MetS phenotype explained by the hippurate diet score

after taking into account all covariates (age, sex, family relatedness and BMI [for triglycerides]).

(3) The proportion of the variance in each MetS phenotype explained by the hippurate diet score after taking into account all covariates as in (2) and adjusting for the applicable variable.

Appendix F. Chapter 7 Appendices

Table 1. Nutrient profile and linear trends of the visceral fat mass diet score according to score tertile

Tertile 1		Tertile 2	Tertile 3	Trend	
Nutrient	Mean (SD)	Mean (SD)	Mean (SD)	Beta (SE)	Р
Energy (kcal)	1892.1 (510.9)	1804.2 (522.9)	1870.4 (552.2)	-15.61 (14.56)	NS
Fat (g/d)	61.5 (10.4)	68.6 (9.5)	72.4 (10.3)	5.53 (0.28)	1.62x10 ⁻⁷⁵
Saturated FAs (g/d)	21.3 (5.4)	24.5 (5.2)	26.6 (5.1)	2.65 (0.14)	1.90x10 ⁻⁶⁷
MUFAs (g/d) 19.7 (3.7)		22.7 (3.4) 24.7 (3.8)		2.54 (0.1)	2.33x10 ⁻¹¹³
PUFAs (g/d)	15.2 (4.4)	15.9 (4.4)	15.7 (4.5)	0.3 (0.12)	1.21x10 ⁻²
trans-FAs (g/d)	1.3 (0.6)	1.6 (0.6)	1.8 (0.6)	0.25 (0.02)	2.55x10 ⁻⁵¹
Cholesterol (mg/d)	202.5 (69.1)	234.8 (74.2)	251.5 (78.2)	24.98 (2.06)	4.04x10 ⁻³²
Protein (g/d)	79.5 (12.2)	81 (12.3)	79.6 (12.5)	0.15 (0.34)	NS
Carbohydrate (g/d)	248.1 (29.6)	229 (29.1)	217.2 (31.1)	-15.66 (0.84)	1.13x10 ⁻⁶⁹
Starch (g/d)	107.7 (26.3)	107.9 (25.1)	111.2 (26.1)	1.65 (0.71)	2.05x10 ⁻²
Total sugars (g/d)	137.5 (28.1)	118.3 (24.4)	103.4 (24.1)	-17.22 (0.71)	5.34x10 ⁻¹⁰⁸
Glucose (g/d)	28 (9.2)	22.4 (7.8)	18 (6.5)	-5.05 (0.22)	8.20x10 ⁻¹⁰¹
Fructose (g/d)	33.7 (11.4)	26 (9.2)	19.9 (7.5)	-6.99 (0.26)	1.29x10 ⁻¹²⁵
Sucrose (g/d)	48 (14.1)	44.2 (14.4)	42.7 (15.6)	-2.73 (0.41)	6.34x10 ⁻¹¹
Maltose (g/d)	3.3 (1.5)	3 (1.6)	3.3 (1.8)	-0.02 (0.05)	NS
Lactose (g/d)	18.7 (9.9)	18.3 (8.9)	17 (9.8)	-0.82 (0.26)	1.82x10 ⁻³
NSP (g/d)	22.9 (5.5)	19.6 (4.5)	16.4 (4.2)	-3.22 (0.13)	2.27x10 ⁻¹¹⁰
Alcohol (g/d)	8.4 (11.1)	8.8 (11.3)	10.9 (14.9)	1.24 (0.38)	9.45x10 ⁻⁴
Water (g/d)	2754.6 (662)	2593.7 (598.9)	2478.5 (598.1)	-139.45 (16.71)	1.80x10 ⁻¹⁶
Sodium (mg/d)	2308.6 (498.6)	2262.8 (471.7)	2162.2 (453.5)	-71.5 (12.87)	3.33x10 ⁻⁸
Potassium (mg/d)	4080.4 (624.4)	3833 (569.2)	3550.3 (524.2)	-263.97 (15.56)	1.48x10 ⁻⁵⁸
Chloride (mg/d)	3705.3 (764.9)	3598.8 (728.6)	3411.9 (694.5)	-144.23 (19.7)	4.30x10 ⁻¹³
Calcium (mg/d)	1095.7 (286.9)	1039.4 (260.8)	949.3 (264.1)	-72.17 (7.31)	3.18x10 ⁻²²
Magnesium (mg/d)	369.1 (52.3)	336.3 (49.2)	296.7 (42.4)	-36.01 (1.27)	3.72x10 ⁻¹³⁸
Phosphorous (mg/d)	1534.9 (218.8)	1474.9 (202.7)	1357 (211.8)	-87.14 (5.81)	4.40x10 ⁻⁴⁷
Iron (mg/d)	13.5 (3.1)	12.5 (2.4)	11.5 (2.3)	-0.99 (0.07)	2.43x10 ⁻³⁹
Copper (mg/d)	1.6 (0.4)	1.6 (0.5)	1.5 (0.5)	-0.07 (0.01)	1.90x10 ⁻⁸
Zinc (mg/d)	10.1 (1.5)	10.1 (1.6)	9.8 (1.7) -0.13 (0.04)		2.60x10 ⁻³
Manganese (mg/d)	4.4 (1.1)	4 (1)	3.6 (0.9)	-0.43 (0.03)	9.54x10 ⁻⁵²
lodine (ug/d)	212.9 (73)	207 (68.7)	196.5 (68.7)	-8.03 (1.89)	2.25x10 ⁻⁵
Retinol (ug/d)	407.1 (403.2)	533.9 (586.6)	617.4 (772.6)	106.49 (16.78)	3.05x10 ⁻¹⁰
Carotene (ug/d)	6060.1 (4193.3)	5399.6 (3002.5)	4729.8 (2742.7)	-664.87 (94.64)	3.41x10 ⁻¹²
Vitamin D (ug/d)	2.5 (1.1)	2.5 (1)	2.4 (0.9)	-0.05 (0.03)	5.36x10 ⁻²
Vitamin E (mg/d)	11.7 (3.4)	10.9 (2.9) 9.9 (2.8)		-0.92 (0.08)	1.02x10 ⁻²⁶
Thiamin (mg/d)	1.8 (0.4)	1.7 (0.3) 1.6 (0.3)		-0.12 (0.01)	3.38x10 ⁻³¹
Riboflavin (mg/d)	2.3 (0.7)	2.2 (0.6)	2.1 (0.6)	-0.09 (0.02)	2.03x10 ⁻⁷
Niacin (mg/d)	21.7 (5)	21.1 (4.5) 20.5 (4.5)		-0.59 (0.13)	4.09x10 ⁻⁶
Tryptophan (mg/d)	16.6 (2.5)	16.9 (2.5) 16.6 (2.5)		0.02 (0.07)	NS
Vitamin B6 (mg/d)	2.6 (0.6)	2.5 (0.5)	2.3 (0.5)	-0.15 (0.01)	2.63x10 ⁻²³
Vitamin B12 (mg/d)	5.8 (2.2)	6.3 (2.5)	6.4 (2.9)	0.31 (0.07)	6.65x10 ⁻⁶
Folate (ug/d)	416 (116)	378.8 (103.5)	341.8 (91.7)	-37.1 (2.79)	5.95x10 ⁻³⁸
Pantothenate (mg/d)	6.1 (1.6)	6 (2.8)	5.6 (1.3)	-0.23 (0.04)	1.43x10 ⁻⁸
Biotin (ug/d)	46.9 (10.1)	45.5 (9.4)	42.3 (9.5)	-2.27 (0.26)	1.50x10 ⁻¹⁷
Vitamin C (mg/d)	198.9 (81.5)	164.1 (66.9)	125.8 (50.8)	-36.44 (1.82)	8.20x10 ⁻⁷⁸

NS= not significant: P>0.05. FA: fatty acid, MUFAS: monounsaturated fatty acids, PUFAS: polyunsaturated fatty acids, NSP: non-starch polysaccharides

Tertile 1: score 0-6; tertile 2: score 7-9; tertile 3: scores 10-15. Linear trend determined by using the tertile of the VFM diet score as a predictor of the energy-adjusted nutrient intake.

Table	2. Associations	between viscera	al fat mass (VFM) and the VFM	diet score,	Shannon
Index	and top microbi	ome and metabo	lite associations	s in the MZ disc	cordant twir	n sample

				Regression		
		Low-VFM twins	High-VFM twins	VFM		
Variable	N pairs	Mean(SE)	Mean(SE)	Beta(SE)	Р	R ²
VFM diet score	80	-0.141(0.122)	-0.047(0.115)	0.281(0.091)	0.002	0.057
Alpha- hydroxyisovalerate	80	-0.239(0.120)	0.064(0.111)	0.141(0.094)	0.136	0.014
Bilirubin (Z,Z)	68	0.093(0.102)	-0.110(0.134)	-0.187(0.102)	0.068	0.023
Butyrylcarnitine	80	-0.168(0.126)	0.096(0.124)	0.199(0.087)	0.023	0.032
Hippurate	80	0.062(0.112)	-0.162(0.114)	-0.297(0.095)	0.002	0.058
Shannon Index	27	-0.019(0.182)	-0.268(0.224)	-0.287(0.176)	0.110	0.048
<i>Bifidobacterium</i> OTU (4426298)	27	0.140(0.718)	0.175(1.098)	-0.064(0.197)	0.749	0.002
Eubacterium dolichum	27	-0.108(0.155)	-0.013(0.171)	0.131(0.227)	0.566	0.006

Notes: A linear regression was conducted using the VFM diet score, Shannon Index and top microbiome and metabolite associations to predict VFM in the MZ discordant (1 SD apart in VFM) twin sample. Variables standardized to have mean=0, SD=1.

Appendix G. Publications throughout my PhD

<u>Pallister, T.</u>, Spector, T.D. Food: a new form of personalised (gut microbiome) medicine for chronic diseases? Sep 2016 In: *Journal of the Royal Society of Medicine*. 109, 9, p. 331-336.

*I wrote this review in a joint effort with Professor Tim Spector.

<u>Pallister, T.</u>, Haller, T., Thorand, B., Altmaier, E., Cassidy, A., Martin, T., Jennings, A., Mohney, R.P., Gieger, C., MacGregor, A., Kastenmüller, G., Metspalu, A., Spector, T.D., Menni, C. **Metabolites of milk intake: A metabolomic approach in UK twins with findings replicated in two European cohorts.** 28 Jul 2016 In: *European Journal of Nutrition.* [Epub ahead of print]

*Work for this publication appears in Chapter 4.

<u>Pallister, T.</u>, Jennings, A., Mohney, R. P., Yarand, D., Mangino, M., Cassidy, A., MacGregor, A., Spector, T. D. & Menni, C. **Characterizing blood metabolomics profiles associated with self-reported food intakes in female twins.** 29 Jun 2016 In: *PL o S One.* 11, 6, e0158568

*Work for this publication appears in Chapter 4.

Pallister, T., Sharafi, M., Lachance, G., Pirastu, N., Mohney, R. P., MacGregor, A., Feskens, E. J. M., Duffy, V., Spector, T. D. & Menni, C. Food Preference Patterns in a UK Twin Cohort. Dec 2015 In: *Twin Research and Human Genetics*. 18, 6, p. 793-805

*I created an online food preference questionnaire based on Dr Valerie Duffy's questionnaire. I used this questionnaire for validating the food metabolite scores in Chapter 5.

<u>Pallister, T.</u>, Spector, T. D. & Menni, C. Twin studies advance the understanding of gene-environment interplay in human nutrigenomics. Dec 2014 In: *Nutrition Research Reviews*. 27, 2, p. 242-251

*I wrote this literature review which appears in Chapter 1.