Inter-individual differences in response to dietary intervention: integrating omics platforms towards personalised dietary recommendations

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Technologic advances now make it possible to collect large amounts of genetic, epigenetic, metabolomic and gut microbiome data. These data have the potential to transform approaches towards nutrition counselling by allowing us to recognise and embrace the metabolic, physiologic and genetic differences among individuals. The ultimate goal is to be able to integrate these multi-dimensional data so as to characterise the health status and disease risk of an individual and to provide personalised recommendations to maximise health. To this end, accurate and predictive systems-based measures of health are needed that incorporate molecular signatures of genes, transcripts, proteins, metabolites and microbes. Although we are making progress within each of these omics arenas, we have yet to integrate effectively multiple sources of biologic data so as to provide comprehensive phenotypic profiles. Observational studies have provided some insights into associative interactions between genetic or phenotypic variation and diet and their impact on health; however, very few human experimental studies have addressed these relationships. Dietary interventions that test prescribed diets in well-characterised study populations and that monitor system-wide responses (ideally using several omics platforms) are needed to make correlation–causation connections and to characterise phenotypes under controlled conditions. Given the growth in our knowledge, there is the potential to develop personalised dietary recommendations. However, developing these recommendations assumes that an improved understanding of the phenotypic complexities of individuals and their responses to the complexities of their diets will lead to a sustainable, effective approach to promote health and prevent disease – therein lies our challenge.

Abbreviation: GST, glutathione S-transferase.

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Nutrition, as a science, has a long tradition of determining the nutrient requirements of heterogeneous populations eating a wide variety of diets and of providing dietary recommendations for health. This has typically involved simplifying the inherent complexity into manageable recommendations in the form of dietary guidance for the purpose of preventing disease in a population. Despite the application of biostatistical approaches with the goal to be as inclusive of the population as possible, there are limitations due to assumptions that metabolic organisational structure is uniform among individuals and that direct cause–effect relationships exist. In reality, the large number of functional redundancies and adaptive mechanisms that provide for homoeostasis make evaluating the complexities and nuances challenging.

The concept of a ‘nutritional phenotype’, i.e. an integrated set of genetic, proteomic, metabolomic, functional and behavioural factors that, when measured, could provide the basis for assessment of human nutritional status, was introduced several years ago by Ziesel et al. It was proposed as a way to integrate the effects of diet on disease/wellness and provide a quantitative indication of the paths by which genes and environment exert their effects on health. The concept provides a good base from which to begin to establish approaches to personalised dietary recommendations; however, several questions need to be addressed. These include, but are not necessarily limited to: What data will we need on an individual in order to personalise dietary recommendations? How can we use controlled feeding studies and other dietary interventions to generate a nutritional phenotypic framework? How can we most effectively integrate omics data so as to be able to apply them towards personalised nutrition?

What data will we need on an individual in order to personalise dietary recommendations?

Numerous factors contribute to variation in nutritional requirements and responses to diet, including sex, stage of life cycle, disease, physical activity level, genetic background, gut microbial community and environmental exposures. Several of these are already considered in the construction of personalised nutritional recommendations; for example, sex, age, adiposity and activity level are routinely used in determining nutrient requirements in healthy individuals and understanding the contributions of disease state to nutritional requirements is a hallmark of therapeutic nutrition. To date, the more complex factors such as genomics, host microbial community structure and environmental exposures are often not included in the equation.

Genetic polymorphisms are well-recognised sources of variation in human response to some aspects of diet, including taste preference, food tolerance, nutrient absorption, transport and metabolism, and effects at target tissues. Typically, in past studies, one particular genetic variant has been considered in relation to intake of one particular nutrient. For example, two polymorphisms in the MTHFR gene (C677T and A1298C) are associated with reduced methylenetetrahydrofolate reductase activity and higher homocysteine concentrations. Carriers of these polymorphisms are at higher risk of CVD; thus sufficient intake of folate is particularly important. Other examples include iron overload and haemochromatosis, copper malabsorption and Menkes disease, and glucose-6-phosphate dehydrogenase and consumption of fava beans, high in pro-oxidant glycosides (favism; reviewed in ). Further, genomics may contribute to phenotypic differences in health behaviour and modify response to interventions designed to change health behaviours.

Several genome-wide association studies have evaluated the association between multiple SNP and metabolomics profiles. In a sample of 284 men, Gieger et al. integrated genome-wide association study data with serum metabolomics-based quantitation of 363 metabolites. They reported associations of frequent SNP with differences in the metabolite homoeostasis, explaining up to 12% of the variance. Using ratios of certain metabolite concentrations as a proxy for enzymatic activity, up to 28% of the variance can be explained (P-values 10–16–10–21). Four variants in genes coding for enzymes (FADS1, LIPC, SCAD and MCAD) were identified where a corresponding metabolic phenotype (metatype) clearly matched the biochemical pathways in which these enzymes are active.

More recently, Suhre et al. conducted an analysis of genotype-dependent metabolic phenotypes using a genome-wide association study with non-targeted metabolomics in a sample of 1768 individuals. They identified thirty-seven genetic loci associated with blood metabolite concentrations, of which twenty-five showed effect sizes that accounted for 10–60% difference in metabolite levels per allele copy. These results provided functional insights into disease-related associations that have been reported in previous studies, including those for cardiovascular and renal disorders, type 2 diabetes, cancer, gout, venous thromboembolism and Crohn’s disease.

The human gut microbial community also shapes host exposure to dietary constituents by modulating absorption, storage and energy harvest from the diet. It is a large, complex ecosystem, with the number of different species of bacteria estimated to range from 300 and 1000 and the majority of the species diversity distributed between the phyla Firmicutes and Bacteroidetes. There is high inter-individual variation in the composition of communities, mostly at the species level, whereas the distribution of bacterial functional genes is less varied. This functional redundancy is a hallmark of a stable symbiosis in which many different species carry out the same functional role.

Recent studies suggest that individuals can be clustered into distinct groups based on their gut microbiome composition and functional metabolism. The underlying metabolism of the dominant bacteria that define these groups is the degradation of plant polymers (e.g. dietary fibre) via different metabolic pathways; long-term dietary habits have been associated with these groupings. Through the metabolism of dietary constituents, the gut microbiome can influence the magnitude and flux of metabolites to which the host is exposed and some of the variations in what have been identified as genotype-dependent metabolic phenotypes actually may be due to the composition and activity of the gut microbiome.
Indeed, of the genotype-dependent metabolic phenotypes
identified by Suhre et al.\(^{16}\), an altered microbiome has
been associated with CVD\(^{13}\), type 2 diabetes\(^{14}\), some
cancers\(^{15,16}\) and Crohn’s disease\(^{17}\). However, the rela-
tionships between the gut microbiome, diet and metabolic
phenotypes need to be addressed in rigorous experimental
settings using approaches that integrate metabolomics, host
genomics and the gut microbiome.

How can we use controlled feeding studies and other
dietary interventions to develop phenotype profiles?

Controlled feeding studies in healthy human subjects have
been used for over a century to establish the quantitative
requirements and confirm essentiality of nutrients in
human subjects. Typically, these studies had small sample
sizes, were intensively controlled, and often focused on
restriction and re-feeding of specific nutrients or nutrient
sources. They were used to evaluate the acute effects of
food deprivation, show experimentally the effects of diet-
ary restrictions on development of deficiency diseases,
establish specific amino acid requirements and describe
vitamin metabolism\(^{18}\). Consequently, they were crucial in
determining recommended daily dietary allowances. Con-
trolled interventions and defined background diets have
also been useful for testing response to varying doses of a
dietary constituent\(^{19}\) and for testing and monitoring bio-
markers of disease susceptibility and dietary exposure\(^{20}\).
More recently, dietary interventions have been used to test
the effects of particular dietary patterns\(^{21}\) and to test
genotype–phenotype interactions\(^{22}\).

Controlled feeding studies, particularly with randomised
crossover designs where each person serves as their own
control, are a useful venue in which to test genotype–diet
interactions as well as genotype–phenotype associations. In
the latter case, the relationship between genotype and
phenotype can sometimes be better characterised on the
background of the same dietary exposures (i.e. a controlled
diet)\(^{22}\). Participant screening protocols for recruitment
into controlled feeding studies also can be set up to enrich
a priori for particular genotypes or phenotypes so as to
provide more equal distributions of sample sizes in sub-
groups, particularly if the prevalence of a particular variant
is low, and to increase statistical power to compare these
subgroups.

Controlled feeding studies also provide a useful
approach in which to characterise host-gut microbial
interactions and to determine gut microbial community
response to diet. In the context of controlled dietary inter-
ventions, gut bacterial community composition has been
shown to differ significantly when participants consume
different diets\(^{23,24}\), although the overall response of the
gut bacterial community is often unique for each individu-
al\(^{24,25}\). Most studies have tested effects of fermentable
complex carbohydrates (e.g. dietary fibres, resistant starch;
Table 1). Network analysis of the gut microbial community
reveals niche specialisation based on a metabolic inter-
connection between different bacteria that are often spe-
cialised in one enzymatic transformation in the pathway of
dietary metabolism\(^{10,26–28}\). The type of carbohydrate
ingested often influences the prevalence of certain groups
of gut bacteria and the subsequent composition of the
microbial metabolic end products to which the host is
exposed (e.g. SCFA; Table 1). Differences in gut microbial
metabolism of various phytochemicals also contribute to
gut bacterial metabolic phenotypes that influence dietary
exposures\(^{29}\). Being able to test for the effects of these
phenotypes in the context of nutrition interventions is
important, since some subgroups may be more responsive
to the intervention than others. For example, Niculescu
et al.\(^{30}\) reported differential lymphocyte gene expression
by bacterial metabolic phenotype in postmenopausal
women receiving an isoflavone supplement; a greater
increase in oestrogen-responsive genes was observed in
women who carried the bacteria capable of converting the
soya isoflavone daidzein to equol.

‘Omics’ – transcriptomics, proteomics and metabo-
lomics – approaches have been hypothesised to revolutio-
nise our understanding of the interactions of the various
systems that are often studied in isolation and have the
potential to revolutionise many aspects of our study of
nutrition and health promotion. Despite the excitement,
at this stage, the technologies still require rigorous evaluation
and validation. Controlled feeding studies are a useful
approach in which to validate and test the robustness of
these omics approaches with the goal of ultimately being
able to use them to evaluate the effects of totality of diet
on totality of response in human subjects. In addition, they
provide important details on the behaviour of proteins,
transcripts and metabolites under controlled conditions.

Several studies have used the construct of controlled
feeding interventions to test effects of diet on omics
measures (Table 2). The majority of these have utilised
metabolomics to characterise response to phytochemical-
containing foods (fruits, vegetables, tea, nuts) compared
with a control in healthy individuals. Many of the meta-
bolites identified typically correspond to dietary bio-
markers of the intervention foods consumed (e.g. proline
betaine after consumption of citrus fruits). Although many
studies also yield a handful of endogenous metabolites that
differ in abundance between the interventions, these com-
ounds are often generally reported as differences in
metabolite profiles owing to a lack of adequate pathway
analysis tools. Thus, it is often unclear whether differences
in metabolite profiles are indicative of perturbations in
specific pathways or molecular targets in response to the
dietary intervention, or are unrelated compounds identified
by chance. Some investigators have explored pathways
manually. For example, Solanky et al.\(^{31}\) found that soya
consumption was associated with osmolyte fluctuations
and differences in energy metabolism. Work in our
laboratory (DH May, SL Navarro, I Ruczinski et al., un-
published results) suggests potential differences in energy
utilisation from glucose to fat between a diet devoid of
fruits and vegetables compared with a diet high in cruci-
fers, citrus and soya. These examples provide provocative
views of other mechanisms through which plant foods may
promote health; however, even with manual analyses, the
interpretation is still broad, speculative and incomplete.

Other investigations have employed alternative omics
technologies to study response to diet and have evaluated
### Table 1. Summary of human dietary randomised crossover intervention studies of response of the gut microbiome to diet

<table>
<thead>
<tr>
<th>Dietary intervention in normal weight individuals</th>
<th>Sample size/population</th>
<th>Dietary intervention/dose of food agent</th>
<th>Treatment duration</th>
<th>Platform/biological sample/key outcomes</th>
</tr>
</thead>
</table>
| Hooda et al.  
([56]) | n 20 M; healthy adults | Randomised controlled crossover, no fibre, polydextrose (21 g/d); soluble maize fibre (21 g/d) | 21 d | Pyrosequencing of V4 region of 16S rRNA gene/faeces. PCA showed gut microbial community shifts with fibre interventions.  |
| Ross et al.  
([57]) | n 17; 11 F, 6 M | Randomised controlled crossover, WG(150 g/d) v. refined grain | 2-week | Bacterial enumeration using FISH; *C. leptum* group increased in WG diet along with stool frequency.  |
| Costabile et al.  
([58]) | n 31; 16 F, 15 M | Randomised controlled crossover, 48 g/d breakfast cereal of either 100% WG (11.8 g DF/100 g; chosen after a pre-screening for bifidogenicity) or WB (27 g DF/100 g) ad-lib diet; energy composition was not the same; WG contained higher content of non-sugar carbohydrate  | two 3-week periods | Bacterial group enumeration using FISH showed changes in several groups with both WG and WB diets and differences between diets. No change in faecal SCFA. Increase in fasting plasma furic acid with WB  |
| Finley et al.  
([59]) | n 40 pre-metabolic syndrome; n 40 controls; 20 F, 20 M | 1/2 cup (130 g) of pinto bean puree v. chicken–noodle soup  | 4-week equilibration and 12-week intervention | Faecal bacterial species enumeration using FISH. Breath methane measured. No effect of bean consumption, except *E. limosum* levels decreased by 50%  |
| Smith et al.  
([60]) | n 18 M | Single-blind randomised crossover, self-managed with addition of seven experimental foods (bread, muffin, brownie, choc milk drink, muesli, pasta, mashed potatoes) with or without LKF; LKF diet provided 17–30 g additional fibre/d; mean fibre intake was 23 g/d on control and 45 g/d on LKF; LKF diet was sig. lower in starch  | 28-d ;3-d pooled faecal collection at end of each period | FISH of 16S rRNA genes/faeces with probes for total bacteria and specific groups. No difference in total bacteria, but changes in certain groups in response to treatment.  |
| Johnson et al.  
([61]) | n 38 M | Single-blind crossover, same as Smith et al. [60] aforementioned | 28-d; 3-d pooled faecal collection at end of each period | SCFA and bacterial enzyme activity in faeces. LKF altered bowel function parameters and decreased faecal pH. Faecal SCFA increased and β-glucuronidase activity decreased. No difference in faecal ammonia concentration  |
| Tuchy et al.  
([62]) | n 31; 17F, 14M | 20 g FOS +10 g PHGG/d v. placebo | crossover, 3-week feeding periods; two consecutive d stool sample mix | FISH of 16S rRNA genes. Increase in *Bifidobacterium* spp., but no difference in total bacteria between diets or other spp. enumerated. No change in faecal pH  |
| Hylla et al.  
([63]) | n 12; 5 F, 7 M | Randomised controlled crossover, high v. low resistant starch: amylomaize starch in bread, pasta, cake and biscuits (Hylon VII, Natl Starch) | 4-week | Changes in breath hydrogen and several faecal parameters associated with bacterial activity (e.g. pH, certain SCFA, β-glucosidase, secondary bile acids)  |

### Dietary intervention in overweight/obese individuals

<table>
<thead>
<tr>
<th>Sample size/population</th>
<th>Dietary intervention/dose of food agent</th>
<th>Treatment duration</th>
<th>Platform/biological sample/key outcomes</th>
</tr>
</thead>
</table>
| Weickert et al.  
([64]) | n 69; 43 F, 26 M | Randomised controlled crossover, high cereal fibre 43 g/d, moderately high cereal/fibre/protein diets, 23% energy as protein, cereal fibre 26 g/d | 18 weeks | FISH/Flow cytometry of faecal bacteria. No effect of diet on bacterial groups  |
| Russell et al.  
([65]) | n 17 M; obese | Randomised controlled crossover. Maintenance diet (85 g protein, 116 g fat and 360 g carbohydrate/d) HPMC (139 g protein, 82 g fat and 181 g carbohydrate/d) HPLC (137 g protein, 143 g fat and 22 g carbohydrate/d) | 7 d maintenance followed by 14-d intervention | FISH of 16S rRNA genes/GC-MS analysis of faecal water content/faeces. HPMC and HPLC diets resulted in increased proportions of branched-chain fatty acids and concentrations of phenylacetic acid and N-nitroso compounds. HPLC diet decreased proportion of butyrate in faecal SCFA concentrations, concomitant with reduction in Roseburia/Eubacterium rectale bacteria, and reduced fibre-derived, antioxidant phenolic acids  |
Table 1 (Continued)

<table>
<thead>
<tr>
<th>Dietary intervention/dose of food agent</th>
<th>Sample size/population</th>
<th>Dietary intervention, reference, and year</th>
<th>Treatment duration</th>
<th>Platform/biological sample/key outcomes</th>
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<tr>
<td>3-d maintenance/medium (HPMC; 24 g/d, 35%) or low carb (HPLC; 24 g/d, 4%)</td>
<td>n = 19 M, n = 12; obese adults</td>
<td>n = 19 M</td>
<td>3-d maintenance</td>
<td>Bacterial enumeration using FISH showed shifts in certain groups of bacteria. Faecal SCFA and ammonia lower during experimental periods than maintenance and proportions of some SCFA differed by diet. Pyrosequencing of 16S rRNA gene. Ratio of Bacteroidetes to Firmicutes decreased over time associated with weight loss. No difference between diets</td>
</tr>
<tr>
<td>4-week experimental diet</td>
<td>n = 36</td>
<td>n = 36</td>
<td>4-week experimental diet</td>
<td>Reduced fat about 30% of calories from fat or 25% of calories from carbohydrate; fibre was 10–15 g/d. Weight loss. No difference between diets</td>
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<tr>
<td>1 year</td>
<td>Reduced fat about 30% of calories from fat or 25% of calories from carbohydrate; fibre was 10–15 g/d. Weight loss. No difference between diets</td>
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</table>

DF, dietary fibre; F, female; FISH, fluorescent in-situ hybridisation of 16S rRNA genes; FOS, fructo-oligosaccharide; HPMC, high-protein and moderate-carbohydrate; HPLC, diet and a high-protein and low-carbohydrate; LKF, lupin kernel fibre; M, male; PCA, principal components analysis; PHGG, partially hydrolysed guar gum; WB, wheat bran; WG, whole grain.

How can we most effectively integrate omics data so as to be able to apply them towards personalised nutrition?

Given that cellular functions are carried out via orchestrated activities of multiplex components of biological systems, data from different omics platforms can shed light on cellular activities at different levels. Methods that integrate omics data from different molecular profiling studies, e.g. data from transcriptomics, proteomics or metabolomics studies, have the potential to provide new insight into how different components of biological systems interact with each other and form the basis of an individual’s health. Here, we provide an overview of available methods of data integration from multiple omics platforms, provide examples of each of different approaches, and discuss their advantages and limitations.

Current methods for integrative analysis of omics data from multiple data platforms can be broadly grouped into three categories. The first class of models, which we refer to as concordance analysis methods, studies concordance/correlation between two omics datasets, e.g. comparing the gene expression levels and proteomics datasets on the same other endpoints beyond differences in metabolite profiles. Brauer et al. (32), interrogated the proteome in response to 2 weeks of a diet high in cruciferous vegetables, and assessed whether response differed by glutathione S-transferase (GST) M1 genotype. GST enzymes metabolise a variety of exogenous compounds, including isothiocyanates from cruciferous vegetables, and the GST M1 variants resulting in a complete lack of gene product are common (22). Twenty-four distinct peaks were associated with cruciferous vegetable consumption compared with a fruit- and vegetable-free diet, two of which were identified that changed in a GST M1-genotype-dependent manner. Another study provides an example of a novel use of omics to link metabolic phenotypes with dietary preferences. Taking a targeted approach, Rezzi et al. (33) used lipidomics to determine metabolites associated with chocolate ‘desiring’ or chocolate ‘indifferent’ preferences among individuals consuming 50 g/d chocolate or bread as a placebo. Heinzmann et al. (34) used metabolomics to study the stability of phenotypic response to diet through sequential dietary challenges. They found that inter-individual differences were often greater than differences within an individual in response to dietary modulation, providing evidence that individuals each have a unique metabolic phenotype. Moreover, intra-individual differences between consecutive dietary challenges were linked to differences in excretion of microbial co-metabolites suggesting flexibility in gut microbiome function in response to dietary modulation. As the authors point out, these differences illustrate the importance of assessing response to diet in the context of a crossover rather than parallel study design in order to move towards personalised nutrition. As a whole, these controlled feeding studies illustrate the potential for omics technology in characterising individual nutritional phenotypes, but make evident the challenges (i.e. compound identification, pathway analysis) that still exist.
### Table 2. Summary of human dietary intervention studies using metabolomic and proteomic platforms

<table>
<thead>
<tr>
<th>Dietary intervention, reference and year</th>
<th>Sample size/population</th>
<th>Dietary intervention/dose of food agent</th>
<th>Treatment duration</th>
<th>Platform/biological sample/key outcomes</th>
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</thead>
<tbody>
<tr>
<td><strong>Fruits and vegetables</strong></td>
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<tr>
<td>May D, Navarro SL, Ruzcinski I et al.</td>
<td>n 10; 5 F, 5 M; healthy adults</td>
<td>Randomised controlled crossover, mixture of cruciferous vegetables, citrus fruits and soya (F&amp;V) compared to fruit and vegetable-free diet (basal); 5 g/kg BW</td>
<td>2 weeks</td>
<td>Metabolomics/8 h fasting urine; more abundant in the F&amp;V: markers of dietary intervention (e.g. crucifers, citrus and soya metabolites), fatty acids and niacin; more abundant in basal: riboflavin, several acylcarnitines and amino acid metabolites; differences in energy utilisation between diet treatments</td>
</tr>
<tr>
<td>Van Dorsten et al. (66)</td>
<td>n 58 (29 in each treatment arm); 25 F, 33 M hypertensive adults</td>
<td>Randomised double-blind placebo-controlled double-crossover, capsules containing a polyphenol rich mix of either red wine and red grape juice extracts (800 mg) or only red grape extract (800 mg), and placebo</td>
<td>4 weeks</td>
<td>Metabolomics/urine; 18 phenolic acids elevated after either polyphenol treatment including syringic acid, 3- and 4-hydroxyhippuric acid and 4-hydrohippuric acid and 4-hydroxymandelic acid</td>
</tr>
<tr>
<td>Brauer et al. (32)</td>
<td>n 36; 17 F, 19 M; n 42; 17 F, 25 M; healthy adults recruited based on GSTM1 genotype (present or null)</td>
<td>2 separate randomised controlled crossover trials of mixed vegetables: (1) 436 g cruciferous; 90 g allium; and 270 g apiaceous; (2) 7 g/kg BW cruciferous; 14 g/kg BW cruciferous; 7 g/kg BW cruciferous + 4 g/kg BW apiaceous; both compared to fruit and vegetable-free diet (basal)</td>
<td>6 d; 2 weeks</td>
<td>Proteomics/8 h fasting serum; 24 distinct peaks associated with cruciferous vegetables; 20 associated with GSTM1 genotype; joint study analysis showed 6 peaks changed in genotype-dependent manner; two identified as TTR and ZAG</td>
</tr>
<tr>
<td>Heinzmann et al. (67)</td>
<td>n 8; 7 F, 1 M healthy adults</td>
<td>Standardised mixed fruit meal (apple, orange, grapes and grapefruit; no dosage provided)</td>
<td>3 d</td>
<td>Metabolomics/urine; excretion of proline betaine, tartaric acid, hippuric acid and benzoic acid was increased compared to baseline</td>
</tr>
<tr>
<td>Walsh et al. (68)</td>
<td>n 21; 12 F, 9 M healthy adults</td>
<td>Non-controlled crossover, 2 d habitual diet; 2 d low phytochemical diet and 2 d high phytochemical diet (100 ml x 4 apple, carrot and strawberry drinks)</td>
<td>2 d</td>
<td>Metabolomics/fasting urine; higher excretion of hippurate and lower excretion of creatinine and methylhistidine discriminated the high phytochemical and habitual diets from the low phytochemical diet</td>
</tr>
<tr>
<td><strong>Other plant foods</strong></td>
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<tr>
<td>Tulipani et al. (69)</td>
<td>n 42; adults with metabolic syndrome</td>
<td>Randomised parallel intervention, mixed nuts, 30 g/d and control</td>
<td>12 weeks</td>
<td>Metabolomics/urine; 20 potential markers of nut intake including fatty acid, phase II, microbiobly-derived phenolic, and serotonin metabolites</td>
</tr>
<tr>
<td>Llorach et al. (70)</td>
<td>n 24; healthy adults</td>
<td>Randomised blind placebo-controlled, encapsulated almond skin extract, 3-5 g</td>
<td>Single dose of ten capsules</td>
<td>Metabolomics/urine; 34 metabolites of almond skin including flavonoids, hydroxyphenylvalerolactone, 4-hydroxy-5-(phenyl):valeric acid, hydroxyphenylpropionic acid, hydroxyphenylacetic acid and other phenolic acid conjugates</td>
</tr>
<tr>
<td>van Dorsten et al. (71)</td>
<td>n 17 M; healthy adults</td>
<td>Randomised crossover, black tea (6 g/d), green tea (6 g/d) or caffeine (control)</td>
<td>2 d</td>
<td>Metabolomics/urine; green and black tea increased urinary excretion of hippuric acid and 1.3-dihydroxyphenyl-2-O-sulfate; greater increase in several citric acid cycle intermediates with green tea</td>
</tr>
<tr>
<td>Solanky et al. (31)</td>
<td>n 9 F; pre-menopausal</td>
<td>Controlled, miso (50 g/d; n 6) or soya protein (60 g/d; n 3)</td>
<td>4 weeks</td>
<td>Metabolomics/urine; increased TMAO, methylamine, dimethyl amine, choline, creatine, glutamine (soya only) and glutamate (soya only), and decreased creatinine, hippurate, benzoate, citrate (miso only) and lactate (miso only)</td>
</tr>
<tr>
<td>Dietary intervention, reference and year</td>
<td>Sample size/population</td>
<td>Dietary intervention/dose of food agent</td>
<td>Treatment duration</td>
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<tr>
<td><strong>Miscellaneous</strong></td>
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<tr>
<td>Rasmussen et al.(72)</td>
<td>77; 44 F, 33 M; overweight, adults</td>
<td>Randomised, high (23–28% of energy; n 42) or low (10–15% of energy; n 35) protein diet</td>
<td>6 months</td>
<td>Metabolomics/urinary; creatine increased with a high protein diet; citric acid increased with the low protein diet</td>
</tr>
<tr>
<td>Moazzami et al.(73)</td>
<td>17 M; prostate cancer patients</td>
<td>Randomised controlled crossover, whole grain rye, rye bran and refined white wheat product, control (485 g/d)</td>
<td>6 weeks</td>
<td>Metabolomics/fasting plasma; metabolites increased after rye bran included 3-hydroxybutyric acid, acetone, betaine, N,N-dimethylglycine and dimethyl sulfone</td>
</tr>
<tr>
<td>Heinzmann et al.(34)</td>
<td>7; 6 F, 1 M healthy adults</td>
<td>Controlled, various dietary challenges including mixed fruit (apple, orange, grapes and grapefruit), fish, wine and grapes, beef and fish</td>
<td>7 d</td>
<td>Metabolomics/urine; inter-individual metabolic differences were greater than effects of any single dietary challenge; differences to dietary challenges differed between individuals</td>
</tr>
<tr>
<td>Zivkovic et al.(74)</td>
<td>3; 1 F, 2 M; healthy adults</td>
<td>Standardised test beverages, 40% kcal needs; 230 g lactose-free milk, 227 g low-fat yogurt, 30 g 100% whey protein powder, 118 g banana and 22 g flax seed oil</td>
<td>Three single test beverages</td>
<td>Targeted lipidomics/plasma; serum fatty acid differences were greater among individuals than within; three metabolites discriminated individuals in ApoB fraction; TAG16 : n7, TAG18 : 2n6 and phosphatidylcholine18 : 3n3</td>
</tr>
<tr>
<td>Llorach et al.(75)</td>
<td>10; 5 F, 5 M; Healthy adults</td>
<td>Randomised crossover, 40 g cocoa powder with water or 250 ml milk or 250 ml milk alone</td>
<td>Single test beverages</td>
<td>Metabolomics/urine; 27 cocoa-phytochemical metabolites identified after both cocoa-containing beverages</td>
</tr>
<tr>
<td>Bertram et al.(76)</td>
<td>28; 8-year-old boys</td>
<td>Randomised, 53 g/d protein from low-fat milk or low-fat meat</td>
<td>7 d</td>
<td>Metabolomics/urine and serum; urinary hippurate excretion was decreased with milk; urinary creatinine, histidine and urea was increased with meat</td>
</tr>
<tr>
<td>Rezzi et al.(33)</td>
<td>75 M; healthy adults</td>
<td>Controlled crossover, chocolate (50 g) or bread (placebo)</td>
<td>Single feedings</td>
<td>Lipidomics/urine and plasma; metabolic phenotypes associated with chocolate desiring or chocolate indifferent preferences</td>
</tr>
<tr>
<td>Stella et al.(77)</td>
<td>12 M; healthy adults</td>
<td>Randomised crossover, vegetarian (420 g/d), low-meat (60 g/d) or high-meat (420 g/d)</td>
<td>15 d</td>
<td>Metabolomics/urine; urinary excretion of carnitine, creatinine, taurine, TMAO, methyhistidine was increased with high-meat; p-hydroxyphenylacetate increased after vegetarian diet</td>
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BW, body weight; F, female; GSTM1, glutathione S-transferase M1; M, male; SMCSO, S-methyl-L-cysteine sulfoxide; TMAO, trimethylamine-N-oxide; TTR, transthyretin; ZAG, zinc α-2-glycoprotein.
set of subjects. The objective of such an approach is to identify genes/proteins/metabolites with an orchestrated activity in a given biological setting. To this end, methods of multivariate analysis, including different variations of principal component analysis, partial least squares, self-organising maps, as well as methods of network visualisation and analysis, have been used to assess the associations among multiple datasets. For instance, Hirai et al.\textsuperscript{[35]} applied principal component analysis as well as self-organising maps to discover relationships between transcriptome and metabolome in Arabidopsis. In another study, Hirai et al.\textsuperscript{[36]} analysed the network of gene-to-gene and gene-to-metabolite associations. More recently, Cao et al.\textsuperscript{[37]} proposed a sparse partial least squares procedure for comparative analysis of data from two omics platforms and applied their method to data from cDNA and Affymetrix chips in NCI60 cancer cell lines.

Concordance analysis methods provide interesting information about components of biological systems that interact with each other in a given setting. Moreover, such analyses can lend themselves to better classificatory models based on a combination of biomarkers from different platforms. However, these approaches often provide limited new insight into the underlying biological mechanisms as omics data from different platforms often show low levels of correlation due to complex mappings of genes to proteins and metabolites, and various post-transcriptional events\textsuperscript{[38]}. Further, the underlying assumption in the majority of these methods is that omics measurements are obtained on the same set of individuals, or more formally, share a common dimension. Van Deun et al.\textsuperscript{[39]} reviewed these different approaches for analysis of multiple omics data, in the setting where the datasets share a common set of features.

The second class of integrative models, which we refer to as sequential integration, includes methods that incorporate multiple sets of omics data in order to discover new biomarkers or delineate biological mechanisms of complex phenotypes. It uses multiple omics datasets, in a sequential manner, to further narrow down, or expand, the set of biomarkers. Sequential integration methods can exploit different methods of data analysis, from simple differential expression analysis to gene-set enrichment analysis or analysis of networks. In examples of such an approach, Puturi et al.\textsuperscript{[40]} first identified the set of differentially active metabolites, and then used meta genomic data to identify pathways associated with prostate cancer progression. In another study, Puturi et al.\textsuperscript{[41]} coupled this approach with a concordance analysis based on metabolomics flux measurements to delineate pathways and biomarkers associated with bladder cancer. More recently, Imieliński et al.\textsuperscript{[42]} used gene-set enrichment analysis coupled with the knowledge of biological networks and compared two sequential approaches, called ‘gene-centric’ and ‘protein-centric,’ in a study of molecular bases of breast cancer. In each of these approaches, the authors first evaluated the enrichment of biological pathways based on one source of data (transcriptomic or proteomic) and then filtered the set of identified pathways based on the second source of data. The authors also compared the results of these methods with a concordance-based approach, where the pathways were identified based on gene and protein pairs that demonstrated orchestrated levels of activity.

Sequential integration methods offer an opportunity to gain new insight based on multiple sources of omics data. Moreover, these methods do not require the omics measurements to be necessarily observed for the same set of individuals. Finally, unlike methods of concordance analysis, which cannot be directly extended to analysis of more than two sets of omics data, sequential integration methods offer the flexibility of analysing multiple omics datasets. However, the power of these methods is clearly limited by the ability of the omics data chosen for the first stage of analysis to capture important biological mechanisms. As the study by Imieliński et al.\textsuperscript{[42]} indicates, the results of the analysis can vary depending on the omics platform used for the first stage of analysis. This sensitivity to the order of analysis can potentially hinder the applications of sequential integration methods, and additional studies are needed to determine whether data-driven criteria can be developed to assess the optimal order of analysis in these methods.

The final group of omics integration techniques, which we refer to as concurrent integration methods, includes emerging approaches that attempt to address some of the shortcomings of the afore-mentioned two sets of approaches. Similar to sequential integration methods, concurrent integration methods try to exploit the information content of multiple sets of omics data. However, these methods often include measures of activity of biological pathways, or their components, based on multiple omics data. This is often achieved by defining a combined score for the activity of each pathway based on activities of its members measured by different omics datasets. Poisson et al.\textsuperscript{[43]} compared the performance a number of methods for combining data from multiple omics platforms, by considering different summary measures defined based on individual test statistics, with methods based on a single omics data source and show that the integrative approaches can improve the power of the analysis. In a recent study, Jauhiainen et al.\textsuperscript{[44]} proposed a multivariate approach, using a mixed linear model, to assess the association of transcriptomics and metabolomics measurements with cancer progression. The proposed model requires measurements to be observed on the same set of samples, but offers the potential for discovering novel biological mechanisms, as well as biomarker identification. On the other hand, Shojaie et al. (A Shojaie, K Panzit, N Puturi et al., unpublished results) propose a network-based method, based on the NetGSA method\textsuperscript{[45]}, for integrating multiple sources of omics data, which can be applied to data from different samples. This procedure does not lend itself directly to selection of biomarkers, and follow-up analyses are needed to determine which components of the selected pathways should be used as biomarkers.

Concurrent integrative methods have also been proposed for gaining insight into biological mechanisms in the cell. An example of such an approach includes the proposal of Shojaie et al. (A Shojaie, A Jauhiainen, M Kallitsis and G Michailidis, unpublished results) to integrate perturbation screens and steady-state gene expression profiles for discovering causal genetic regulatory mechanisms. In this
study, the authors compare their proposed integrative approach with state-of-the-art methods based on a single source of omics data, and show superior estimates of regulatory networks can be obtained that by combining multiple omics data. Table 3 summarises the different classes of integration methods.

Novel biomedical technologies continue to improve the quality of the omics data, as well as to reduce the cost of obtaining such data. In nutrition studies, biological experiments now generate multiple sources of omics data including transcriptomic, proteomic, metabolomic and gut microbial community measurements. The main challenge is now integrating such measurements in a systematic way, in order to provide a holistic view of biological systems. As more and more measurements become available, the complexity of the analysis, i.e. the number of variables in statistical models, increases. This poses additional challenges for design of trials, and necessitates the use of advanced statistical models appropriate for analysis of high-dimensional problems. A potential solution for this challenge is to incorporate available biological knowledge, including information on biological pathways and genetic, protein interaction and metabolic networks. Incorporating biological information can both reduce the dimensionality of the problem, and also improve the power and reproducibility of analysis methods.

**A way forward to personalised nutrition**

There is still a lot of effort needed to establish a robust health phenotype framework on which to develop personalised dietary recommendations. The improving omics technologies and the ability to integrate various omics platforms in a systematic fashion will facilitate providing a holistic view of cellular functions related to healthy phenotypes; however, the characterisation of the contribution of diet to the biochemical and metabolic parameters associated with healthy phenotypes would benefit from systematic evaluation under controlled conditions in well-described groups of individuals. Controlled human feeding studies are a useful experimental setting in which to conduct this work. Nonetheless, these types of studies are expensive and funding multiple, new large-scale dietary interventions that capture a variety of dietary patterns and intakes is likely to be prohibitive.

An efficient and effective way to develop some of the necessary omics databases under experimental conditions may be to take a collaborative approach, leveraging existing samples from previously conducted human interventions. Stored samples from controlled feeding studies are stashed away in freezers around the globe and in many cases are well characterised and ideal for further omic analysis. Statistical techniques for integrating multiple omics data from a common platform but different study populations, i.e. meta-analysis techniques, already exist; they improve statistical power by integrating samples from different study populations in which to conduct this work. Nonetheless, these types of studies are expensive and funding multiple, new large-scale dietary interventions that capture a variety of dietary patterns and intakes is likely to be prohibitive.

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### Table 3. Summary of methods for integrative analysis of multiple omics datasets

<table>
<thead>
<tr>
<th>Integration approach</th>
<th>Reference</th>
<th>Methodology/tools</th>
<th>Omics data</th>
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<tr>
<td>(2) Sequential integration</td>
<td>Putluri <em>et al.</em>[40]</td>
<td>DE, OCM</td>
<td>Metabolomics, meta-genomics in prostate cancer Metabolomics abundance and flux data, meta-genomics in bladder cancer Transcriptomics, proteomics in breast cancer</td>
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DE, differential analysis; GSEA, gene-set enrichment analysis; CA, correlation analysis; PCA, principal component analysis; SOM, self-organising maps; PLS, partial least squares; OCM, Oncomine concept mapping; NetGSA, network-based gene-set analysis.
of health phenotypes, several factors need to be considered if personalised nutrition is to move towards being a part of routine health practice. Adherence to dietary recommendations for chronic disease prevention at the population level, such as those of national and international associations (e.g. US Department of Agriculture, World Cancer Research Fund, American Heart Association) is associated with lower risk of chronic disease; for example, greater adherence to the 2005 US Dietary Guidelines was inversely associated with risk of CHD, stroke, diabetes and total cancer.\(^{[55]}\) In theory, tailored recommendations may be more closely associated with risk of CHD, stroke, diabetes and total cancer.\(^{[55]}\) In practice, finding individualised approaches that facilitate and maintain desired dietary behaviour on the heels of a personalised diet prescription for health will likely remain an ongoing challenge for nutrition practitioners.

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