

Perspective: Metabotyping—A Potential Personalized Nutrition Strategy for Precision Prevention of Cardiometabolic Disease

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ABSTRACT

Diet is an important, modifiable lifestyle factor of cardiometabolic disease risk, and an improved diet can delay or even prevent the onset of disease. Recent evidence suggests that individuals could benefit from diets adapted to their genotype and phenotype: that is, personalized nutrition. A novel strategy is to tailor diets for groups of individuals according to their metabolic phenotypes (metabotypes). Randomized controlled trials evaluating metabotype-specific responses and nonresponses are urgently needed to bridge the current gap of knowledge with regard to the efficacy of personalized strategies in nutrition. In this Perspective, we discuss the concept of metabotyping, review the current literature on metabotyping in the context of cardiometabolic disease prevention, and suggest potential strategies for metabotype-based nutritional advice for future work. We also discuss potential determinants of metabotypes, including gut microbiota, and highlight the use of metabolomics to define effective markers for cardiometabolic disease-related metabotypes. Moreover, we hypothesize that people at high risk for cardiometabolic diseases have distinct metabotypes and that individuals grouped into specific metabotypes may respond differently to the same diet, which is being tested in a project of the Joint Programming Initiative: A Healthy Diet for a Healthy Life. *Adv Nutr* 2020;11:524–532.

Keywords: metabotyping, personalized nutrition, precision nutrition, targeted nutrition, cardiometabolic diseases, gut microbiota, metabolomics

Introduction

Diet is among the most important modifiable lifestyle factors contributing to cardiometabolic disease risk. Changes in a diet can delay or even prevent the onset of disease. In the current work, we define cardiometabolic disease as various disease entities that are either consequences of or late stages of the metabolic syndrome. We therefore include metabolic syndrome, Type 2 diabetes, and cardiovascular disease endpoints in this definition. Prevention is considered to be the most sustainable and cost-effective way to manage chronic diseases (1). While national and international dietary guidelines serve to promote health and prevent disease from a population perspective, data from Europe, the United States,

and Australia have clearly shown that these guidelines are poorly adhered to (2–6). To combat obesity and related conditions, some individuals may need to follow personalized nutrition strategies as a complement to the general population-based advice. Such recommendations have been suggested in the most recent advisory report for the Dietary Guidelines for Americans, based on recent randomized controlled trials showing that personalized interventions could lead to greater weight loss than non-personalized strategies (7). In addition to health status being improved as a result of improved biological effects when personalizing the diet, there is also evidence that people are more open to health-promoting information when it is personalized and

when the individuals recognize themselves as being highly susceptible to preventable diseases (8, 9). However, it remains to be elucidated to what extent personalized nutrition has advantages over generalized strategies and whether it is feasible at a large scale (10–12).

Personalized nutrition at the individual level requires not only the comprehensive collection of information, which is both costly and demanding, but also models that are capable of accurately generating personalized advice for the individual. A more feasible approach may be to personalize diets at the group level. Over the last 10–15 years, individuals have been grouped according to genetic set up, for example based on genetic variants that are associated with disease risk (13). However, the benefits to public health have been limited. More recent studies have suggested that individuals may be grouped according to unique metabolic responses to foods and dietary changes (14, 15). In fact, what constitutes a healthy diet may differ across individuals as a consequence of a number of factors beyond genetics, including health status, medication, and also the gut microbiota (16–18). Grouping individuals based on similarities in their metabolic phenotype—that is, metabotypes—is a novel concept, and different definitions have been used (19–23). The underlying idea behind metabotyping is to identify metabolic phenotypes based on factors such as diet, anthropometric measures, clinical parameters, metabolomics data, and the gut microbiota. An optimal diet can then be tailored to fit each metabotype specifically (Figure 1). In this perspective article, we discuss the current literature reporting on metabotypes in the context of cardiometabolic disease and how metabotyping may be used as a nutritional strategy for the improved prevention of chronic disease.

Current Status of Knowledge

Definitions and strategies for personalized nutrition

There is currently no overall, generally agreed-upon definition of personalized nutrition or precision nutrition. As in other scientific fields under development, concepts have

been used often without uniform definitions. In the area of personalized nutrition, concepts adapted to individuals or groups of individuals have been described: Ordovas and colleagues (10) defined “stratified and tailored nutrition” as approaches attempting to tailor diets to groups of individuals sharing similar characteristics, “personalized nutrition” as an attempt to deliver nutritional interventions or advice suited to each individual, and “precision nutrition” as the most ambitious definition, which implies a quantitative understanding of the individual, their food intake, and their phenotype (including health) to offer the nutritional interventions or advice of the greatest benefits to the individual. Furthermore, Gonzalez and Betts (24) integrated dynamics—that is, a time dimension—in their definition of precision nutrition, aiming to tailor nutrition to best meet the varying demands and requirements throughout a day, week, season, or lifespan. Most recently, the North American Branch of the International Life Sciences Institute convened a multidisciplinary panel and defined personalized nutrition as using “individual-specific information, founded in evidence-based science, to promote dietary behavior change that may result in measurable health benefits.” In addition, they suggested 10 guiding principles for personalized nutrition approaches (25). Here, we use personalized nutrition and precision nutrition interchangeably, and define it as the most appropriate diet to maximize health benefits tailored to the individual or a group of individuals.

It has been suggested that targeted nutritional advice can be personalized on 3 levels: on the basis of the current diet (Level 1); on the diet and phenotype, consisting of anthropometric and biochemical/clinical measurements (Level 2); or diet, phenotype, and genotype (Level 3) (26–29). Some evidence suggests that strategies that include genetic information have greater potential than strategies based on the background diet and health phenotype alone for improving health through personalized nutrition (28, 30–32). However, omics techniques other than genomics may also benefit personalized nutrition, such as analyses of the microbiome (metagenomics) and the metabolome (metabolomics), but these techniques have not been included in the investigation of different personalization levels. Although such omics techniques provide phenotype information, represented at Level 2, they hold immense potential compared to the traditional phenotypic measurements, as they provide a functional readout of a phenotype at a much more detailed level. Metabolomics can, to some degree, reflect food and nutrient consumption, as well as provide mechanistic evidence of the specific metabolic pathways impacted by diet (33, 34), and the gut microbiota phenotype has been shown to modulate physiological responses to diet (18).

In a landmark article from 2015, Zeevi et al. (16) showed benefits when adapting diets based on factors underlying inter-individual differences in post-prandial glucose response. In brief, the post-prandial glucose response could be accurately predicted using a tree-based machine learning algorithm, combining data on gut microbiota, blood markers,

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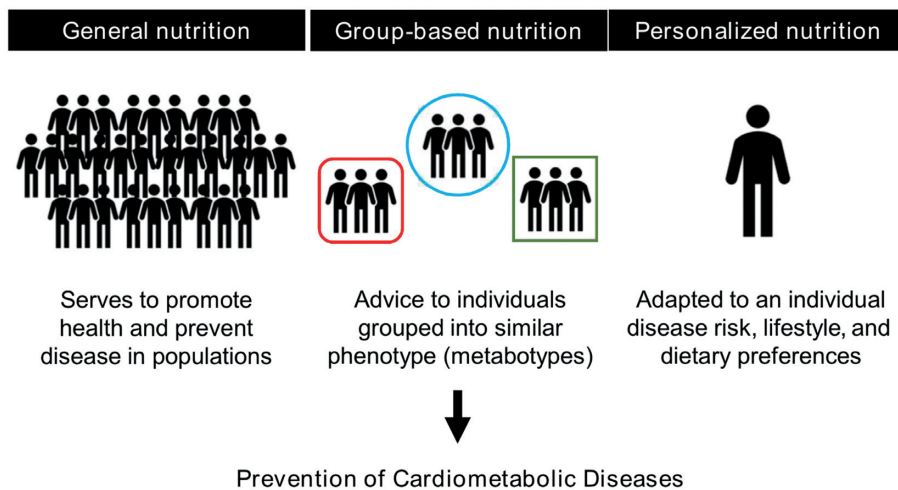


FIGURE 1 Metabotyping and group-based nutrition in the context of the conventional population-based guidelines and personalized nutrition.

body composition, diet, and exercise. Next, the authors were able to perform personalized interventions with “good” and “bad” diets, capable of affecting post-prandial glucose levels in a favorable or non-favorable direction. Importantly, what constituted a “bad diet” for some participants resembled the “good diet” for other participants, and vice versa. This study thus showcased the potential to use phenotypic measures in combination with machine-learning algorithms on biological big data for precision nutrition at the individual level. The approach was recently validated in an American cohort of non-diabetic subjects (35). Similar phenotyping strategies can be used to identify metatypes and, thus, personalize nutrition on a group level.

Metabotyping based on anthropometric and clinical/biochemical parameters

Metabotyping, where individuals were clustered into metabolic phenotypes based on anthropometric, biochemical, and clinical cardiometabolic measures, has been suggested as a potential nutritional strategy (21, 36–42). Riedl et al. (43) showed that 3 metatypes, derived from BMI and 33 biochemical parameters (e.g., blood lipids, glucose, hormones, and liver enzymes) using k-means clustering, differed in cardiometabolic disease prevalence and 7-year disease incidence, thus illustrating the ability to detect high- and low-risk groups. Krishnan et al. (44) detected unique, post-prandial leptin and glucose responses for 3 distinct groups of overweight, healthy women in a cross-over study on high- versus low-glycemic index meals, using a principal component analysis. Although a firm link between post-prandial glucose response and long-term health outcomes has yet to be established among healthy subjects, differences in post-prandial metabolism are considered reflective of the metabolic health status of individuals and serve as a proxy for high-risk groups of cardiometabolic disease (45). Interestingly, when reanalyzing

data from a weight-loss trial, Ritz and colleagues (46) found that metabotyping based on fasting glucose and insulin could predict weight loss from a healthy, Nordic diet rich in dietary fiber, versus a control diet (46). They also concluded that these basic measures, although seemingly of clinical relevance, likely lacked details on metabolic regulation, and that more complex measurements could provide better personalized nutrition advice. Moreover, O’Donovan and colleagues (21) demonstrated a communication strategy for large-scale, metatype-based personalized nutrition: a decision-tree approach providing each participant with pre-defined dietary messages based on their metatype, in combination with their BMI, waist circumference, and blood pressure. The 3 metatypes had been identified based on 4 blood-based, clinical measures: blood glucose, triacylglycerols, total cholesterol, and high-density lipoprotein cholesterol. Compared to the personalized advice delivered by a dietitian, the decision tree generated comparable recommendations.

Gut microbiota as a potential determinant of metatype

In recent years, the immense role of the gut microbiota in cardiometabolic disease has been highlighted, and this has been an area of intensive research (47–49). The impact of the gut microbiota on human health could partly be mediated through the ability of the gut microbiota to metabolize dietary compounds into new metabolites that impact disease risk. For example, the bacterial metabolism of dietary L-carnitine into the metabolites trimethylamine and trimethylamine N-oxide presents a mechanistic link of the relationship between red meat consumption and cardiovascular disease risk (50), and short-chain fatty acids provide a functional link between gut microbiota, dietary fiber intake, and several diet-related diseases (51–53).

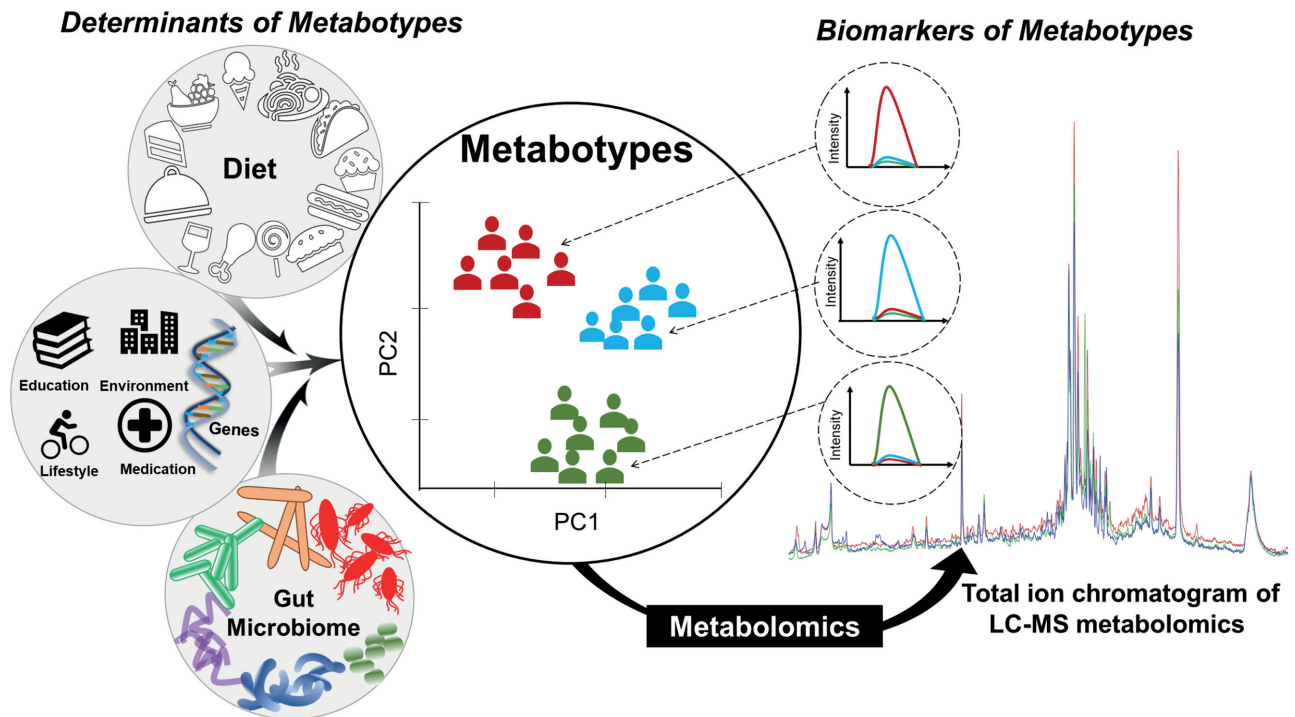


FIGURE 2 The graphic illustration of potential determinants of metatypes and a metabolomics-based strategy for the identification of biomarkers of metatypes by using untargeted LC-MS metabolomics. Individuals are grouped into metatypes based on potential determinants and their interactions, using certain statistical approaches, such as PC analysis. PC, principal component.

Interestingly, the diet–microbiota interactions have been shown to vary amongst individuals (54) and appear to be a determinant of the post-prandial response, as previously discussed (16). Moreover, even when a diet seemingly induces more systematic effects on the phenotype, a large part of the residual variability is associated with the microbiota in terms of composition and functionality (55). Several researchers have observed that individuals with similarities in their microbiota composition also share similarities in their phenotypic response to exposures. For instance, Romo-Vaquero et al. (56) observed that the *Coriobacteriaceae* could play a central role in urolithin production metatypes (i.e., metatypes that reflect the unique patterns of microbiota-derived metabolites from ellagitannin-rich foods), which in turn were associated with blood cholesterol levels and cardiometabolic health implications. The grouping of individuals by microbiota phenotype (enterotype) (57, 58) is a concept similar to metotyping, with the important distinction that enterotyping is based on the microbial composition, with potential impact on diet–health interactions, whilst the metatype partly reflects the activity of the microbiota in conjunction with dietary exposures and other phenotypic traits. The enterotype may thus be an important determinant of the metatype, and studies investigating enterotyping in the context of diets can, therefore, provide unique and valuable insights into how to integrate metotyping into precision nutrition (Figure 2).

For example, Kovatcheva-Datchary et al. (18) demonstrated the potential to predict response to diet on the

enterotype level by showing that the prevalence of *Prevotella* determined individuals’ glycemic response to barley-based bread in healthy adults. The contribution of the enterotype was further shown by fecal transplants from study participants to germ-free mice. Improved glucose regulation and a higher capacity to digest dietary fiber were observed only in mice of the enterotype rich in *Prevotella*. Similarly, Korem and colleagues (17) found low- and high-glucose response groups after subjects had consumed whole-grain sourdough bread and refined wheat bread, and the groups could be accurately predicted based on the baseline gut microbial composition. Their results suggest that although there may be underlying, systematic health effects on a population level for different bread types, that effect could be overshadowed by systematic variabilities between individuals. Furthermore, enterotypes based on the relative abundance of the *Prevotella* and *Bacteroides* genera have been shown to impact the response to dietary weight loss interventions, where individuals with a high *Prevotella*-to-*Bacteroides* ratio lost more weight and body fat in response to high-fiber diets, compared to those with a low bacterial ratio (59).

Taken together, enterotypes may be a key determinant of phenotypes and be especially useful when personalizing diets in terms of dietary fiber and glycemic response. Enterotypes are also considered to influence the metabolisms of other dietary compounds, such as polyphenols, leading to the production of enterotype-specific bacterial metabolites (60–62). However, gut microbial genes, which may be important for metabolic disease phenotypes, can be present across

bacterial phyla and also differ between bacterial strains that otherwise share the same genetic makeup (63). This supports the idea of metabotyping: that is, that the functionality of the gut microbiota, rather than the presence of bacteria per se, may be important for disease prevention.

Metabolites as markers for cardiometabolic disease metabolotypes

Metabolomics may be particularly well suited to discover metabolotypes and the biomarkers thereof, as well as to unravel how the gut microbiome contributes to variations in phenotypic responses to diets (Figure 2). Metabolites have been shown to reflect cardiometabolic disease states and risk factors (64–67), as well as host and gut microbial genetics and lifestyle factors (67–69). Moreover, recent studies have shown that microbiota composition can be effectively mirrored in both the fecal and plasma metabolomes (68).

Very few studies have been published where metabolomics has been applied for metabotyping as a means to set up improved strategies to combat cardiometabolic diseases. These studies have clustered men and women into metabolite-derived groups correlated to disease traits, risk factors, and eating patterns (15, 70–72). Men and women from a large Irish study, for example, were clustered according to their plasma fatty acid profile, creating 4 groups that differed in terms of metabolic syndrome components, anthropometric measures, dietary habits, and demographics (70). In another study, urinary metabolites associated with diabetes were used to divide diabetic and non-diabetic participants into 4 novel groups (71). Differences in plasma glucose levels led the authors to speculate on differences in disease management and the risk of future complications for the 2 metabolotypes with diabetic patients. From a nutritional perspective, Fiamoncini and colleagues (15) identified 2 metabolite-based metabolotypes, and evaluated their respective responses to a dietary intervention aimed at weight loss. In brief, healthy, overweight men and women were clustered into 2 distinct metabolotypes based on post-prandial concentrations of metabolites related to lipid metabolism. Individuals belonging to the different metabolotypes also differed in terms of amino acid and carbohydrate metabolism, post-prandial glucose and insulin levels, liver lipid contents, intra-abdominal fat mass, and eating patterns. When challenged with a 12-week weight loss trial with a reduced caloric intake, only the individuals with the more disease-associated metabolotype showed improvements in glucose and insulin levels at the end of the intervention; thus, the study identified a responsive and a nonresponsive metabolotype (15). In another recent study, Muniandy et al. (72) identified 2 distinct subgroups of monozygotic twin pairs associated with cardiometabolic risk factors, high-density lipoprotein cholesterol, and BMI using metabolomics.

A framework for metabolotype-based nutritional advice

Given the heterogeneity in choices of variables which can be used to cluster into metabolotypes, it is difficult to compare

metabolotypes between studies and populations (40). In fact, obtained metabolotypes should be carefully assessed and validated against underlying research questions and, consequently, clustering variables. Planning the experimental study design also encompasses the choice between baseline (fasting or not fasting) and post-prandial samples, biological samples (blood, urine, or fecal samples), and continuous sampling to evaluate responses over time; these decisions are closely linked to the choice of study endpoints. Choosing statistical approaches for deciphering metabolotypes from the large and complex data sets will be an instrumental part of future metabotyping studies and presents a great challenge at present. Part of the difficulty lies in how to handle and weigh the vastly different types of data. Appropriate data analytical methods will not only dictate how metabolotypes are defined, but also reproducibility across research groups and, subsequently, generalizability across study populations. Although beyond the scope of this perspectives article, we note that arbitrary selection of the number of metabolotypes seems to be a common practice to date; however, there are objective methods that select the ultimate number of clusters, as demonstrated by Riedl et al. (40).

Whilst the discovery of metabolotypes may be complex and require careful consideration, metabotyping should preferably be simple, rapid, and affordable once metabolotypes have been defined and validated. Biomarkers suitable for large-scale applications are needed, and we encourage the consideration of metabolotype biomarker evaluation and validation in future metabotyping studies (Figure 2). Blood biomarkers could be useful, as they can be measured repeatedly and continuously, whilst other biological samples (e.g., fecal samples) may be more difficult to collect at defined time points and close time intervals. Moreover, work is needed to evaluate potential underlying mechanistic links and to identify traits of the metabolic phenotype undetected by measurements of a few traditional risk factors, and it may require the inclusion of other data domains, such as gut microbiota and the entire metabolome. Metabolites present as interesting candidates: they may not only be reflective of metabolotypes and responses but have the advantage of being present in several biological specimens, such as blood, urine, and stool. On the other hand, future work should assess whether grouping individuals based on a small number of anthropometric, clinical, or other easily measured parameters alone might be adequate to identify those diets that would be optimal for specific metabolotypes. This is relevant to make personalized nutrition more feasible and cost-effective in clinical practice.

Performing dietary intervention trials that evaluate metabolotype-specific responses is a fundamental step required to bridge the current gap of knowledge with regard to the efficacy of personalized strategies in nutrition. In an ongoing Joint Programming Initiative, A Healthy Diet for a Healthy Life project, “Diet × gut microbiome-based metabolotypes to determine cardio-metabolic risk and tailor intervention strategies for improved health” (73), we have hypothesized that adults at high risk of developing

cardiometabolic disease will have different responses to fermentable dietary fibers, dependent on their metabolotypes and specific diet–gut microbiota interactions. The idea is to define metabolotypes based on the subjects' metabolomes in the context of the gut bacterial profiles, host genetics, lifestyle factors, anthropometric measures, and dietary profile. We will next test our hypothesis in a separate study population using a randomized controlled cross-over dietary intervention; we will investigate the stability of the metabolotypes throughout the intervention and research whether it is possible to shift an unhealthy metabolotype into a more healthy metabolotype. Although the primary endpoint will be postprandial glucose, we will also evaluate other parameters that are important for cardiometabolic diseases, including blood pressure, blood lipids, and circulating C-reactive protein, as a proxy for systemic inflammation. Our study will allow us to not only investigate a new approach for personalized nutrition, but also to provide estimates of the efficacy of such interventions.

Along with short-term interventions to evaluate the effects of specific dietary interventions across metabolotypes on cardiometabolic risk factors, long-term studies to evaluate adherence to metabolotype-specific diets and their role for hard endpoints are warranted, but difficult to perform. However, metabolotyping may also be used in an observational setting in prospective cohort studies. Such a study setting would allow for the assessment of long-term reproducibility of reported metabolotypes and could be used to deduce potential biomarkers, as well as to investigate their generalizability across populations. Recent studies suggest that distinct metabolotypes (and enterotypes) may be present across populations (21, 74), although the prevalence of a given metabolotype may vary between countries (21). Moreover, metabolite-based metabolotypes have been shown to be stable over several years (75, 76), yet some metabolites have shown higher conservation than others (76). Interestingly, metabolotype instability per se has been correlated with the incidence of cardiovascular disease and all-cause mortality (75).

Practical considerations of implementing metabolotype-based precision nutrition in the health-care setting

Personalized strategies rely on the premise that advice can be delivered to individuals or groups of individuals in an effective manner and that such strategies have been proven to be effective and economically feasible. Studies assessing health economic parameters for different personalized strategies, including metabolotyping, are currently lacking and, therefore, needed. For the delivery of personalized nutrition advice, the ability of health-care professionals to communicate targeted nutritional advice, and their likelihood of doing so, will be instrumental if implementing metabolotyping and precision nutrition in the health-care setting, together with support from policymakers, society, and other actors (8).

Several communication strategies have been employed to deliver personalized dietary advice to consumers, differing

in terms of target group, means of communication, and basis of personalization (77). The previously mentioned work by O'Donovan and colleagues also investigated the delivery of metabolotype-specific dietary advice on a larger scale (21, 41). Individuals' willingness and motivation to undergo metabolotyping and nutritional interventions form another practical aspect that warrants consideration. There is a need to evaluate attitudes and acceptance towards metabolotyping, as well as other strategies in precision nutrition. In the meantime, studies investigating attitudes towards genetic testing as a means to prevent disease can provide some insight (78–80). For example, these studies have suggested that adults at high risk for cardiometabolic diseases may be more motivated to change their health behaviors (78); presenting with disease risk factors was associated with a higher willingness to undergo genetic testing, and perceived risks and family medical history were related to the motivation to exercise, as well as to modify lifestyle factors, such as diet and physical activity (79). Yet, the relatively low proportion of participants (14–33%) who reported improving their health behavior 3–9 months following genetic testing also raises the questions of how to achieve high intervention compliance, lasting changes in health behaviors, and improvements in health parameters. We believe it is realistic to assume that a multitude of factors will impact the outcomes and that the contribution of such factors in specific populations needs to be addressed.

Concluding Remarks

Metabolotyping is a relatively new concept within the area of personalized nutrition. In the current literature, metabolotypes are often defined either based on clinical and anthropometric markers (disease-associated metabolotypes) or the metabolism of certain nutrients and dietary components, such as dietary fiber and polyphenols (diet-associated metabolotypes). We hypothesize that the disease-associated metabolotypes—that is, metabolotypes present in those populations at high risk for cardiometabolic diseases—will affect the response to a specific diet. We further hypothesize that the gut microbiota is a key determinant and a modifier of metabolotypes, in addition to habitual diet, genotype, anthropometric measures, and biochemical and clinical markers. In other words, the interaction between the host and exogenous exposures (e.g., diet, drugs, and gut microbiota) is an important factor regarding dietary response, and identifying such “functional” metabolotypes may offer advantages over “clinical biomarker-based” metabolotypes as well as enterotypes for the optimization of a diet to an individual for the prevention of cardiometabolic disease. However, much more work is needed to investigate whether responses to particular diets and dietary items are indeed metabolotype-specific and whether diets tailored for metabolotypes could lead to health improvements that are clinically meaningful. Ongoing and future studies will hopefully: shed light on whether metabolotype clusters can be deduced in general populations, elucidate potential main determinants and, test whether the response to specific dietary interventions, such as diets high

in fermentable fiber, vary across distinct metabolotypes. If so, finding easily measured biomarkers of metabolotypes is our highest priority, to allow the tailoring of diets for optimal prevention at a large scale.

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