Evidence-based precision nutrition improves clinical outcomes by analyzing human and microbial molecular data with artificial intelligence

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Research

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Abstract

Background

Current dietary recommendations are often generalized, conflicting, and highly subjective, depending on the source biases. This results in confusion, skepticism, and frustration in the general population.

Methods

We have developed an objective, integrated, automated, algorithmic approach to diet and supplement recommendations that is powered by artificial intelligence that analyzes individualized molecular data from the gut microbiome, the human host, and their interactions. This platform enables precise, personalized, and data-driven nutritional recommendations that consist of foods and supplements, based on the individual molecular data, to establish and maintain healthy homeostasis.

Results

We describe the application of our precision nutrition technology platform to populations with depression, anxiety, irritable bowel syndrome (IBS), and type 2 diabetes (T2D). In a blinded interventional study, we provided the study participants with precision nutritional recommendations and observed improvements in clinical outcomes by 36% in severe cases of depression, 40% in severe cases of anxiety, 38% in severe cases of IBS, and more than 30% in the T2D risk score that was validated against clinical measurements of HbA1c.

Conclusion

Our AI-driven precision nutrition program achieved statistically significant improvements in clinical outcomes of depression, anxiety, IBS, and type 2 diabetes. These data support the integration of precision food and supplements into the standard of care for these chronic conditions.

Introduction

There is strong epidemiological and molecular evidence that the human diet contributes significantly to the onset and progression of many chronic diseases and cancers [1][2][3][4][5][6]. Dietary advice, however, has been controversial and conflicting. Scientific publications often contradict one another, with one demonstrating health benefits while another suggesting potential harm for the same foods or diets [7][8][9][10]. As an example, one epidemiological study reports that daily consumption of only 100 mL of sugary drinks, even when the source is fruit juice, raises cancer rates significantly [11]. Other studies report that the consumption of daily fruit or fruit juice can actually prevent cancer [12][13][14]. Dairy consumption has also generated contradictory claims, with some evidence supporting an association...
with a higher incidence of some cancers and chronic diseases, some studies showing no such correlations, while other studies demonstrating significant benefits, and USDA recommending dairy to the Americans [3][15][16][17][18]. Likewise, recent publications provide conflicting data on the effects of saturated fats on cardiovascular and metabolic diseases [15][16][19][20]. Red meats have been touted as healthy sources of critical nutrients, such as vitamin B12 and dietary protein, and even shown to provide health benefits; yet there is strong epidemiological and mechanistic evidence that they may be responsible for increased rates of inflammatory diseases and cancers [1][21][22]. To complicate the landscape further, current nutritional guidelines are influenced by the food industry, which promotes the increased consumption of sugars, dairy, and meat [23][24].

The large body of conflicting nutritional information has manifested in the creation of hundreds of diets, each claiming health benefits. This leaves the consumer confused and frustrated, with little direction or confidence in results. Nutritional science finds itself in this predicament for several reasons. First and foremost, the vast majority of nutritional research doesn't account for the contributions of an individual's gut microbiome, despite a multitude of evidence proving that the impact is significant [25][26][27][28][29]. Secondly, researchers have mistakenly assumed that each food is either good or bad for all humans without understanding that the same food can have very different effects on different people. There is now strong scientific evidence that genetics plays a minor role, and the gut microbiome plays a major role in the effects of nutrition on human physiology [30][31]. This was elegantly demonstrated in a large twins study of metabolic disease parameters [28]. It is clear that the gut microbiome not only influences how food is digested, but also converts many of the molecular ingredients found in foods into beneficial or harmful secondary metabolites that have profound effects on human physiological functions, such as neurotransmitter production, immune system activation and deactivation (inflammation), immune tolerance, and carbohydrate metabolism [32][33][34–36].

Technological advances have made it possible for nutritional science to adopt a paradigm shift. Each food can now be viewed as a container of molecular ingredients (i.e. micronutrients), rather than a homogenous material that is either good or bad for human health. Additionally, the metabolic functions of each person's unique microbiome need to be identified and quantified, in order to "prescribe" specific foods that contain molecular ingredients that will be converted to beneficial (associated with health) secondary metabolites by that person's individual gut microbiome. Microbial functions also need to be quantified so each person can avoid foods that contain molecular ingredients that will be converted by their gut microbiome into harmful (associated with disease) secondary metabolites. In addition to the approaches that utilize known microbial metabolites and their micronutrient precursors, machine-learnt models from large clinical studies need to be applied to each person's gut microbiome analysis to guide the best choice of foods and supplements. Such models have already been developed for selection of foods rich in carbohydrates and can minimize the blood glucose levels, especially in the postprandial compartment [27][26]. Besides the human microbiome, an analysis of human gene expression must be integrated into the systems biology view of the human body to enable the understanding of the network interactions among the dietary molecular ingredients, the microbiome, and human physiology. This understanding is critical because human gene expression, whether modulated by genetic, nutritional,
microbial, or other environmental factors, has a direct connection with the onset and progression of chronic diseases [37, 38][39][40][41].

In this report, we describe an evidence-based approach to improve clinical outcomes using precision diet and supplements that are computed by artificial intelligence algorithms using each individual's molecular data and the self-reported phenotype. This approach is 100% algorithmic and data-driven, and there are no humans involved in making the nutritional recommendations. The molecular data are obtained from stool or a combination of stool and blood samples using highly accurate and reproducible, clinically validated, and clinically licensed (meta)transcriptomic tests [42][43]. These data are used to quantify the activity of microbial metabolic pathways, specific microbial taxa (at the strain level), and human gene expression levels. This information is then converted to precise and personalized diet and supplement recommendations. An overview of this approach is shown in Fig. 1 and described in detail in the sections below. When this approach is used, it can reduce the symptoms of several important and highly prevalent chronic diseases

**Methods**

**Human subjects, ethical considerations, and study design**

The clinical studies described here were approved by a federally-accredited Institutional Review Board (IRB). All samples and metadata were obtained from human subjects at least 18 years old and residing in the USA at the time of participation. All study participants consented to participating in the studies. The study design was non-conventional; while there was no control arm, the subjects were blinded to the fact that they were participating in an interventional study with clinical endpoints. All study participants were recruited into a “wellness study” that asked them to complete a wellness survey. These survey questions were actually clinically validated surveys for depression (PHQ9), anxiety (GAD7), and IBS (Rome IV criteria), or request for their lab results for HbA1c. The clinical studies described here were registered at ClinicalTrials.gov with identifiers NCT04905524 and NCT04905485.

**Metatranscriptomic analyses of stool and transcriptomic analyses of capillary blood samples**

The molecular analyses for the studies reported here focus on sequencing messenger RNAs (mRNAs) isolated from human stool and blood samples. Stool samples were collected and analyzed as previously reported [44]. Briefly, stool samples were collected by the study participants using the Viome commercial kits that included ambient temperature preservation solution and pre-paid return mailers. Stool metatranscriptomic analyses (RNA sequencing, RNAseq) were performed using an automated, clinically-validated laboratory and bioinformatics methods. Results consist of quantitative strain, species, and genus level taxonomic classification of all microorganisms, and quantitative microbial gene and KO (KEGG Ortholog, KEGG = Kyoto Encyclopedia of Genes and Genomes [45]) expression levels. The matching blood samples were collected and analyzed as previously described [42]. Briefly, blood samples
are collected by the study participants using the Viome commercial kits that included ambient
temperature preservation solution and pre-paid return mailers. The kits include lancets and minivettes
that enable easy and accurate collection of small volumes of blood from a finger prick. Transcriptomic
analyses (RNA sequencing, RNaseq) were performed using an automated, clinically-validated laboratory
and bioinformatics methods that require 50 microliters of capillary blood. Test results consist of
quantitative human gene expression data.

All microorganisms that live in the intestines obtain their energy by converting chemical substrates into
products, using metabolic pathways that consist of enzymes. Substrates are typically the molecular
ingredients found in foods, and products are biochemicals usually referred to as secondary metabolites.
Metatranscriptomic analysis of the gut microbiome enables the quantification of thousands of microbial
pathways using the KEGG database. As part of the efficacy trials we describe here, we identified an
average of 377 strains, 363 species, and 102 genera per stool sample, and a total of 2930 strains, 2,007
species, and 451 genera in all stool samples combined. In addition, we have identified an average of
1,916 KOs per stool sample, and 5,467 KOs in all stool samples combined. In blood samples, we
quantified the expression of an average of 11,687 genes per sample and 15,434 genes in all samples
combined.

**Metabolic pathways and functional scores**

We have designed functional scores to quantify certain biological phenomena; for example, leaky gut,
inflammation, gas production, protein fermentation, cellular health, mitochondrial health, etc., that are
relevant to human physiology and healthy homeostasis. Functional scores are weighted functions (\(\text{Score} = C_1 F_1 + C_2 F_2 + \ldots + C_n F_n\), where \(F\) is the feature and \(C\) is its weight) of components from the molecular
data from the gut microbiome and/or blood transcriptome. The components that make up the functional
scores can be taxa, microbial pathways, human pathways, or other functional scores. Functional scores
range from 0 to 100, with 100 representing the highest possible activity. For example, a microbiome-
derived functional score such as Butyrate Production Pathways is calculated as a weighted function
consisting of the expression levels of many known butyrate-associated KOs (Fig. 2) [46]. The expression
level of each KO from the Butyrate Production Pathways score is quantified using the metatranscriptomic
stool test and the score is then computed using the weighted formula [47]. The weights attributed to each
KO within a pathway are determined by a combination of domain knowledge and statistical analyses
obtained from a collection of 200,000 stool and blood samples [48][49]. By quantifying each enzymatic
member (KO) of the butyrate pathways using microbial gene expression data, accurate pathway activities
can be calculated [50].
Table 1: Examples of specific foods and supplements, and reasons for recommendation to consume or avoid.

<table>
<thead>
<tr>
<th>Food/supplement</th>
<th>Reasons to consume</th>
<th>Reasons to avoid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Artichoke</td>
<td>- Low Butyrate Production Pathways functional score, and</td>
<td>- High Gas Production functional score, or</td>
</tr>
<tr>
<td></td>
<td>- Presence of specific microbial strains that convert inulin into butyrate</td>
<td>- Small intestinal bacterial overgrowth (SIBO) symptoms</td>
</tr>
<tr>
<td>Broccoli</td>
<td>- High Inflammatory Activity functional score, or</td>
<td>- High Sulfide Gas Production functional score, and</td>
</tr>
<tr>
<td></td>
<td>- Suspected liver and detox support needed</td>
<td>- High Trimethylamine (TMA) Production functional score</td>
</tr>
<tr>
<td>Trout</td>
<td>High Inflammatory Activity functional score</td>
<td>High Uric Acid Production Pathways functional score</td>
</tr>
<tr>
<td>Nicotinamide Riboside (NR)</td>
<td>- Low Mitochondrial Health functional score, or</td>
<td>- High Cellular Senescence functional score, and</td>
</tr>
<tr>
<td></td>
<td>- Low Mitochondrial Biogenesis Pathways functional score</td>
<td>- High Inflammatory Activity functional score</td>
</tr>
<tr>
<td>Curcumin</td>
<td>- High Inflammatory Activity functional score, or</td>
<td>High Bile Acid Production Pathway functional score</td>
</tr>
<tr>
<td></td>
<td>- High Cellular Stress, or</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- High Immune System Activation</td>
<td></td>
</tr>
</tbody>
</table>

Making personalized nutritional recommendations

Precision nutritional recommendations are computed using the Viome AI Recommendation Engine and are designed to address an individual’s biological patterns at the functional level by boosting beneficial (health-associated) and suppressing harmful (disease-associated) activities with molecular ingredients from foods and supplements (Fig. 1). This approach is built on the concept that on a molecular level, a particular food or supplement may be beneficial for one person, but harmful to a different person. Table 1 shows examples of this concept. The Viome AI Recommendation Engine uses a complex set of algorithms to determine the final food and supplement recommendations. The algorithms are developed from the domain knowledge (publications on microbial and human physiology, food science, clinical trials, etc.), phenotypic information, and extensive clinical studies from which machine-learned models of nutritional modulation of the microbiome and human physiology were developed (e.g. [26]). These algorithms are applied to the functional scores and phenotype for each person whose stool, or stool and blood samples are analyzed. Phenotype is determined from information provided by a participant in the wellness questionnaire, such as symptom assessment, known health conditions, allergies, and medications.
The Viome AI Recommendation Engine considers compounds (molecular ingredients) in foods and supplements that can support the healthy functions of both the gut microbiome and the human. These compounds include specific polysaccharides, polyphenols, vitamins, minerals, amino acids, fatty acids, and many phytochemicals. This approach highlights the concept that a single food is more than simply its macronutrient content and that foods from the same family can have very different molecular compositions. For example, an almond is a source of many phytonutrients and compounds such as kaempferol (flavonoid), naringenin (flavonoid), ferulic acid (phenolic acid), oxalic acid, phytic acid, quercetin, procyanidin B2 and B3, magnesium, phytosterols such as retinol, a-tocopherol, vitamin K, vitamin D, and beta-sitosterol, fatty acids such as oleic acid, linoleic acid, and palmitic acid, and specific amino acids [51]. The decision to recommend a specific compound and its amount depends on the values of multiple functional scores. After considering all inputs, the recommendation engine classifies foods into one of four categories based on the molecular composition of each food. The food categories are superfoods, enjoy foods, minimize foods, and avoid foods, which are consumerized names that correspond to the recommended servings per day for each food.

Personalized supplement recommendations follow the same logic, considering all inputs to identify compounds that are beneficial or harmful to an individual's functional scores and phenotype. Supplements include minerals, vitamins, botanicals or herbs, food extracts, enzymes, phospholipids, amino acids, prebiotics, and probiotics. When considering individual functional scores, supplement ingredients commonly believed to be beneficial may not be recommended. For example, turmeric is a commonly consumed supplement for its anti-inflammatory properties, but has also been shown to increase bile flow [52]. For individuals with a high Bile Acid Metabolism Pathway functional score, turmeric supplementation may be more harmful than beneficial. A high Bile Acid Metabolism Pathway score suggests that the microbial activity of transforming bile salts into bile acids is high. While such biotransformation is part of a balanced gut microbiota and bile acid homeostasis, excessive intestinal bile acids may promote a pro-inflammatory environment and play a role in the development of gastrointestinal diseases [53][54].

The process of categorizing foods and supplements (determining the servings or dose) includes prioritizing scores that need improvement and considering conflicts within the recommendations. An example is shown in Table 2: a low Energy Production Pathway functional score will yield recommendations based on compounds that contribute to the score activity, one of which is alpha-lipoic acid (ALA). Spinach and broccoli are recommended due to the ALA content that is a critical cofactor for mitochondrial energy production enzymes such as pyruvate dehydrogenase (PDH), alpha-ketoglutarate dehydrogenase (alpha-KGDH), and branched-chain ketoacid dehydrogenase (BCKDC) [55]. However, when considering additional score results, broccoli and spinach will be placed on the avoid food list due to broccoli’s glucosinolate content and spinach's oxalate content. Instead, tomatoes and peas are recommended as sources of ALA to support the Energy Production Pathway functional score.
### Example Score Results:

- **High Sulfide Gas Production Pathways functional score**
- **Low Butyrate Production Pathways functional score**
- **Low Oxalate Metabolism Pathways functional score**
- **Low Energy Production Pathways functional score**

### Self-reported information on wellness questionnaire:

- Histamine intolerance
- Pistachio allergy

<table>
<thead>
<tr>
<th>Food</th>
<th>Compounds</th>
<th>Food categories (initial)</th>
<th>Final recommendation</th>
<th>Replacement food/supplement</th>
</tr>
</thead>
</table>
| Spinach   | - Alpha-lipoic acid
- Oxalates | - Superfood, based on low Energy Production Pathways functional score
- Avoid, due to low Oxalate Metabolism Pathways functional score | Avoid                | - Tomato and
- Alpha-lipoic acid supplement |
| Broccoli  | - Alpha-lipoic acid
- Glucosinolates | - Superfood, based on low Energy Production Pathways functional score
- Avoid, due to high Sulfide Gas Production Pathways functional score | Avoid                | - Peas and
- Alpha-lipoic acid supplement |
| Sauerkraut| Probiotics        | - Superfood, based on low Butyrate Production Pathways functional score
- Minimize, due to self-reported histamine intolerance | Minimize             | - Probiotic supplement that includes Lactobacillus acidophilus, Lactobacillus plantarum, Lactobacillus rhamnosus |
| Pistachios| CoQ10             | Superfood for Energy Production Pathways Low
- Avoid due to self-reported pistachio allergy | Avoid                | CoQ10 supplement        |
There are circumstances where a beneficial compound cannot be obtained from food due to an allergy or other health-related issue, or due to the lack of sufficient amounts in food. Personalized supplements help support those gaps in nutrition. In Table 2, pistachios are recommended due to their CoQ10 content. However, if an individual has an allergy to pistachios or exhibits small intestinal bacterial overgrowth (SIBO) symptoms, pistachios will be placed on the avoid or minimize food list. In this situation, CoQ10 can be provided through a supplement as the recommendation engine associates it as beneficial for the same score and/or phenotypic conditions.

### Results

We performed single-arm, blinded, interventional studies where we provided precision recommendations to the study participants and measured the change in clinically validated scores for symptoms of Irritable Bowel Syndrome (IBS), depression, and anxiety.

The primary endpoints for the three conditions were:

- For IBS, the IBS Symptom Severity Score (IBS-SSS, as part of the Rome IV criteria, the Rome Foundation), a clinically validated questionnaire on a scale of 0 through 500, with 300 + indicating severe IBS, 175 to 300 indicating moderate IBS, and 75 to 175 indicating mild IBS.
- For depression, the PHQ9 score, a clinically validated questionnaire of 9 questions that yields a score of 0 through 27, with 15 + indicating moderately severe or severe depression, 10 to 15 indicating moderate depression, and 5 to 10 indicating mild depression [56].
- For anxiety, the GAD7 score, a clinically validated questionnaire of 7 questions that yields a score of 0 through 21, with 15 + indicating severe anxiety, 10 to 15 indicating moderate anxiety, and 5 to 10 indicating mild anxiety [57].

There are several reasons this pilot trial is blinded; the main one is that we did not want the trial participants to know the specific conditions being studied, to minimize the placebo effect. Instead, they were recruited to participate in a wellness study, where we collect survey information. None of the diseases or symptoms were mentioned to the potential participants. We then created a “wellness questionnaire” that included clinically validated surveys for IBS (Rome IV criteria), depression (PHQ9), and anxiety (GAD7), as well as a number of additional general wellness questions.

Each interventional study had a single arm. At time point T1, each participant was given precision nutritional recommendations (diet and supplements) and asked to fill out the wellness questionnaire. At time point T2 each subject was asked to fill out the wellness questionnaire again. There were no additional communications with the participants between the two time points.
Figures 3A, 3B, and 3C show the changes in the interventional study endpoints following the precision recommendations for the three conditions studied. We observe that for all three conditions, IBS, depression, and anxiety, the mean clinical score improvements are highest for the most severe subgroups, at 38% for IBS (mean score reduction = 132.5, t-statistic = 3.60, p = 0.0042), 36% for depression (mean score reduction = 6.05, t-statistic = 6.10, p < 0.0001), and 40% for anxiety (mean score reduction = 7.12, t-statistic = 4.70, p = 0.0022). (We note that the p-value for moderate symptom score of anxiety is 0.0852, which is above the conventional standard of 0.05, but still below 0.1, a threshold considered acceptable in many applications.) All of these improvements are considered clinically significant.

We performed an additional, two-arm interventional study to test the efficacy of our precision nutritional recommendations on people with type 2 diabetes (T2D). The primary endpoint for this study is a T2D risk score built using data from the gut microbiome metatranscriptomic analyses of over 50,000 subjects, and validated against an independent cohort of over 2200 subjects (cite). At time point T1, participants donated their stool sample from which their T2D risk score was calculated, and they were provided their precision nutritional recommendations. At time point T2, these participants donated a second stool sample from which their T2D risk score was calculated. They were also asked how well they adhered to their recommendations, low or high.

To evaluate the effect of adhering to the precision recommendations, we compared the difference in T2D risk scores between the “low adherence” cohort and the “high adherence” cohort, while controlling for other variables. The two cohorts were matched by (a) starting risk score of +/-10 points (i.e., within 20 points) and (b) period of adherence of +/- 60 days (i.e., within a total of 120 days), yielding a cohort size of N = 1456, evaluated over a mean time period of 332 days. Figure 4 shows the difference in the change of the T2D risk score (y axis) between the low-adherent participants (blue box plot) and high-adherent participants (green box plot). We observe a statistically significant risk score improvement for people with high adherence (p = 1.99e-05), with a mean risk score reduction of 7 points [(-30.25) - (-23.21)], which translates to > 30% improvement. This shows that high adherence to our precision nutrition recommendations results in a lower risk of type 2 diabetes.

**Discussion**

We describe a novel technology platform that overcomes some of the major shortcomings of nutritional science and improves clinical outcomes in several highly prevalent chronic diseases [58][59][60]. Traditional nutritional research has been conducted using various available methods. Epidemiological studies have been used to derive nutritional rules using large populations. These have yielded interesting results that have guided certain national-scale dietary recommendations, such as the current USDA guidelines in the USA (Dietary Guidelines for Americans, 2020–2025). Many clinical and pre-clinical studies have also tested the effects of certain foods and diets on various health outcomes. Additionally, many different supplements, including prebiotics and probiotics, and their combinations have been tested for their ability to improve certain symptoms. Most of these approaches have yielded varied results for
different people; that is, some people benefit from the intervention and others do not, or some people are harmed by certain foods or diets, and others are not [61][62][63][64][65][66].

Over the last decade, it has become clear that the gut microbiome plays a crucial role in modulating the human physiology by regulating its immune system, metabolic functions, hormones, and neurotransmitters [67]. It is important to note that the microbiome’s influence on human physiology is exerted via its functions, and not merely the taxonomy [68][69]. For example, the fact that a person’s gut microbiome contains *Faecalibacterium prausnitzii*, which is a well-known butyrate producer, does not mean that it is producing butyrate. However, when it encounters the correct fiber substrate, it will produce this important metabolite [70][71]. *Escherichia coli* is a bacterial species that contains very beneficial members that produce vitamins K and B12, and help human hosts efficiently absorb iron; it also has deadly members, such as the enterohemorrhagic strains [72][73][74]. Therefore, identifying the species of *Escherichia coli* in a gut microbiome is not very informative, given the vastly different functions that members of this taxon can perform.

The key to understanding the effects of the gut microbiome on human physiology and health is to quantify the microbial functions using metatranscriptomic, metaproteomic, or metabolomic approaches. We have chosen metatranscriptomics as the best approach, as it enables quantitative measurements of microbial protein expression that can identify both protein-based and metabolite-based effects on the human physiology, identify molecular ingredients in foods that can modulate microbial functions, and still provide the highest resolution of taxonomic classification (can distinguish strains) that can guide some aspects of the nutritional recommendations. Another essential aspect to understanding the relationship between the human physiology, microbiome and diet is to integrate the human molecular data into the systems biology view of the human body. We use a whole blood transcriptome test that can quantify the expression levels of > 10,000 human genes from capillary (finger prick) blood.

We also want to emphasize that foods should be viewed as containers of molecular ingredients, instead of objects that we can visually recognize as onions, peppers, etc. The reason for this is that each food contains many different molecular ingredients, as exemplified above, that can exert a multitude of health-related effects on the host, either directly or via the microbiome metabolism [75].

**Conclusions**

Here we describe a highly personalized, data-driven, AI recommendation system that uses each person’s molecular data (from stool, or stool and blood) and phenotype (symptoms, medications, etc.) to compute the best diet and supplements for each person. The recommendations are computed in two stages. First, the specific molecular ingredients that should be consumed or avoided are identified, based on which microbial or human functions they support. Next, these molecular ingredients are mapped onto foods that are available in stores. Using this approach, we provided food and supplement recommendations to study participants with signs or symptoms of depression, anxiety, IBS, and type 2 diabetes. We measured clinical outcomes at baseline (prior to delivering the nutritional recommendations) and at follow-up. Our
data show statistically significant improvement in clinical outcomes for all conditions and disease activity levels. This novel precision nutrition approach that uses highly individualized molecular data and machine-learned algorithms should be considered by healthcare professionals and health coaches as an effective supplement to their existing therapeutic strategies.

Limitations Of The Study

Three of the four studies presented here were single arm interventional studies without control arms. However, the participants were blinded to the fact they were in an interventional study, which minimized the placebo effect. We also did not capture the level of adherence to the nutritional recommendations for the studies on IBS, depression, and anxiety, which could significantly affect the observed results (however, we did capture this information for the T2D study, and compare cohorts based on low vs high adherence in Figure 4). It is possible that the observed reductions in the clinical scores would have been higher if the participants with low adherence were excluded from the study. The diet and supplement recommendations were delivered digitally, and no effort was made to either monitor or improve compliance. All of these limitations will be addressed in the future studies.

Declarations

Ethical Approval and Consent to participate

The clinical studies described here were approved by a federally-accredited Institutional Review Board (IRB) in the USA. All samples and metadata were obtained from human subjects at least 18 years old and residing in the USA at the time of participation. All study participants consented to participating in the studies.

Consent for publication

Not applicable.

Availability of data and materials

The data used to generate the figures in this manuscript are available from the corresponding author on reasonable request.

Competing interests

All authors are employees of Viome, Inc., the sponsor of the research described in this manuscript.

Funding

Viome, Inc., is the sponsor of the research described in this manuscript.

Authors' contributions


JC, CH, TL, and GB developed the recommendation algorithms. RT and MV developed the stool and blood tests. RT, GB, and MV developed the study protocols. NS and YC developed machine-learnt algorithms. PM developed digital study algorithms.

Acknowledgements

Not applicable

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2. Endocrinology, Diabetes And Metabolism, AdventHealth, Orlando, FL, USA.

References


**Figures**

*Figure 1*

AI-generated precision nutritional recommendations. Stool and blood samples are collected from participants. The RNAs are extracted, sequenced, and quantified using clinically validated laboratory and bioinformatics methods [44][42]. Microbial genes including KOs are quantified from the RNA sequencing data of each participant. Functional scores are subsequently computed as a weighted function of relevant KOs. Finally, personalized nutrition recommendations are computed using all functional scores and phenotypes.
Figure 2

Microbial butanoate (butyrate) production pathways based on KEGG annotations [46][45].
Figure 3

Efficacy of the nutritional interventions on participants with IBS, depression, or anxiety. A. Results showing the effect of precision nutritional recommendations on subjects with IBS (N=118) over a mean time period of 161 days (time2 - time1), measured using the IBS-SSS score. Bars on the right are mean +/- standard deviation of IBS-SSS score; blue bars represent the baseline (at the study start) and orange bars represent the followup (after the intervention). B. Results showing the effect of precision nutritional
recommendations on subjects with depression (N=143) over a mean time period of 144 days (time2 - time1), measured using the PHQ9 score. Bars on the right are mean +/- standard deviation of PHQ9 score; blue bars represent the baseline (at the study start) and orange bars represent the followup (after the intervention). C. Results showing the effect of precision nutritional recommendations on subjects with anxiety (N=101) over a mean time period of 158 days (time2 - time1), measured using the GAD7 score. Bars on the right are mean +/- standard deviation of GAD7 score; blue bars represent the baseline (at the study start) and orange bars represent the followup (after the intervention). Note that moderate anxiety did not improve as significantly as the others (p=0.09).

<table>
<thead>
<tr>
<th> </th>
<th>Low adherence N=1456</th>
<th>High adherence N=1456</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>46.32 +/- 15.79</td>
<td>49.68 +/- 16.29</td>
</tr>
<tr>
<td>Sex (%female)</td>
<td>62.35</td>
<td>61.82</td>
</tr>
<tr>
<td>Initial risk score</td>
<td>72.31 +/- 25.58</td>
<td>71.84 +/- 25.19</td>
</tr>
<tr>
<td>Period of adherence</td>
<td>261.08 +/- 131.85</td>
<td>255.59 +/- 133.49</td>
</tr>
</tbody>
</table>

**Figure 4**

Efficacy of the nutritional interventions on participants with an elevated T2D risk score. Participants who adhered to precision recommendations (green box) reduced their T2D risk score more than participants who did not (blue box) (p<0.001).