Unraveling the Link between Nutrition and Metabolic Syndrome Risk through In Silico Dietary Interventions

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Abstract

Metabolic Syndrome (MetS) is a cluster of metabolic disorders that substantially increases the risk of chronic metabolic diseases. Diet is known to play a crucial role in the progression of MetS, yet a mechanistic understanding of its impact on MetS risk remains elusive. To address this gap, we conducted a rigorous in silico diet intervention study by leveraging organ-resolved sex-specific whole-body models (WBMs) of metabolism. These models were utilized to computationally evaluate the effect of 12 diverse dietary regimens on key MetS biomarkers—glucose, triacylglycerides (TAG), LDL-C, and HDL-C—and fatty acid beta-oxidation in both males and females. Our analyses elucidated molecular mechanisms underlying the link between conventionally unhealthy diets and elevated MetS risk. Specifically, a typical Unhealthy or Keto diet indicated elevated TAG storage in the adipocytes and increased LDL-C to HDL-C ratios across both genders. Conversely, healthier dietary patterns like the Mediterranean and Vegan diets promoted favorable metabolic profiles, characterized by lower TAG storage and LDL-C to HDL-C ratios. Notably, plant-based (Vegan and Vegetarian) diets induced elevated fatty acid oxidation compared to high-fat regimens, suggesting their potential in mitigating MetS risk. Pronounced gender differences in metabolic responses to diets were also observed in our analyses, highlighting the need for gender-tailored dietary recommendations. Additionally, our study delineated organ-specific contributions to the MetS biomarkers, with the liver and lungs identified as major regulators of blood glucose homeostasis. Overall, this study contributes to a deeper understanding of the intricate interactions between diet and MetS risk.

Introduction

Metabolic Syndrome (MetS) is defined as a cluster of metabolic disorders that significantly increases the likelihood of developing several chronic diseases such as cardiovascular disease and other metabolic disorders, such as Type II Diabetes and Non-Alcoholic Fatty Liver Disease ¹. MetS may involve any combination of metabolic disorders including hypertension (high blood pressure), hyperglycemia (high blood sugar), hyperlipidemia (high blood fat), hypercholesterolemia (high blood LDL-C), polycystic ovary syndrome (PCOS), insulin resistance, and excess body fat around the waist, otherwise known as central obesity ¹. An individual suspected of MetS only needs to display symptoms from three of these metabolic diseases to be eligible for a diagnosis. It is important to note that MetS is not a disease in itself; rather, it is a cluster of interconnected metabolic abnormalities that constitute a pre-morbid condition, increasing the risk of developing other chronic metabolic diseases. MetS has been a growing problem in the Western world and across the globe ². It is now estimated to affect approximately one-third of adults in the United States while its global prevalence ranges from 12.5% to 31.4%, depending on each region's definition of MetS³. The rise in MetS among the general population has significant implications for healthcare costs and the quality of life for affected individuals.

There is growing evidence that diet plays a significant role in developing MetS and is now considered a key modifiable risk factor. Previous studies suggest that diets high in added sugars, refined grains, and unhealthy fats can increase the risk of developing the condition ⁴. On the other hand, diets rich in whole grains, fruits, vegetables, and healthy fats protect against the development of MetS as diets such as a Mediterranean diet, well-balanced, or High Protein diet, are reported to be potentially advantageous for the management of MetS ⁴.

Due to the profound effect of diet on the risk of developing MetS, dietary interventions are seen as an attractive preventive and therapeutic approach for MetS. The common practice in existing dietary intervention studies is to assign one diet to each group of participants to isolate their effects on MetS. Nevertheless, the slowly developing nature of MetS makes it challenging to perform extensive dietary intervention studies to tease out the effect of new diets on MetS development. Additionally, it is practically infeasible to examine the effect of multiple diets on human subjects due to difficulties associated with committing to various new dietary regimens particularly for long-term. Furthermore, since MetS affects many organs and metabolic processes in the body, it is challenging to study and understand the effect of diet on the metabolic function of the many different organs involved in maintaining metabolic homeostasis within the human body.

Computational studies are an intriguing alternative approach in digital medicine to investigate the effect of diet on human metabolism and metabolic disorders such as MetS. For example, a recent study developed a data-driven computational physiological model

describing glucose, lipid, and cholesterol metabolism to investigate the metabolic changes that occur in response to a high-fat, high-sugar diet in mice ⁵. This study showed that this type of diet led to metabolic changes consistent with the development of MetS, including the increased production of fatty acids and inflammation. However, this model was limited only to a small proportion of the metabolism. Furthermore, such computational studies for humans, particularly at genome-scale, are still lacking.

Advances in the development of GEnome-scale Models (GEM) of metabolism have provided a promising route for computationally investigating the effect of diet on human metabolism at genome-scale. GEMs capture all metabolic reactions encoded by the genome of an organism. A global GEM of human metabolism representing the metabolic capability of any human cell without specifying the organ-or cell-type-specific functions, was first reconstructed by Duarte et. al (the Recon1 model) ⁶. This model was then extended by enhancing both topological and functional features, with approximately double the number of reactions and 1.7 times more distinct metabolites (the Recon2 model). The expansion of the Recon models was culminated by the development of the Recon3D model, which contained over 6,000 more reactions than its predecessor, Recon2 ⁷. More recently, another comprehensive GEM of human referred to as Human1 has been developed, which merged the accumulated knowledge from all existing human GEMs into one comprehensive resource ⁸.

Given that these models represent only the global human metabolism, they cannot capture organ-specific metabolic effects. Conversely, while several organ-, tissue-, and cell type-specific GEMs have been reconstructed ⁹, their scope is inherently limited to the metabolic pathways of their specified biological context. A paradigm-shift in this area was the development of two sex-specific organ-resolved whole-body models (WBMs) of human metabolism by Thiele et al ¹⁰. These models integrate GEMs for 20 organs, six sex organs, and six blood cell types into unified sex-specific models for male and female. The models have been parametrized using omics and physiological data and captures both whole-body metabolic processes and the effect of individual organs or cell types on the entire system. The applicability of this model has been demonstrated by integrating it with a dynamic coarse-gained mathematical model of glucose-insulin regulation to study disrupted metabolic processes in Type I Diabetes (T1D) at a whole-body scale ¹¹.

The WBMs offer an unprecedented opportunity to perform extensive in silico dietary intervention studies, taking into consideration organ-specific effects, yet this potential remains untapped in the realm of precision medicine. Here, we sought to use these WBMs to computationally evaluate the effect of a dozen diets on the risk of developing MetS and the role of different organs in this process. We simulated these dietary intakes in silico and evaluated their effects on the contribution of each organ to the serum levels of four key biomarkers of MetS, as well as the activity of the fatty acid oxidation pathway.

Results

In this study, we used the organ-resolved whole-body GEMs of human metabolism for male and female ¹⁰ to computationally investigate the effect of diet on the risk of developing MetS. The male and female WBMs contain 81,094 and 83,521 metabolic reactions, respectively¹⁰. Here, we explored the effect of 12 different diets on the risk of developing MetS. These diets span a wide spectrum of dietary regimens and include an Average American diet, Average European diet, High Protein diet, High Fiber diet, Mediterranean diet, Vegetarian diet, Vegan diet, Gluten Free diet, DACH (Germany, Austria, and Switzerland) diet, Keto diet, a typical Balanced diet, and a typical Unhealthy diet. **Table 1** provides a brief description of each diet. The macronutrient breakdown of each diet is also shown in **Figure 1A**.

To evaluate the effect of these diets on the risk of developing MetS, we mined prior knowledge on the metabolic functionalities and their relevant molecular markers that are associated with the pathophysiology of MetS. This led us to identify four key molecular biomarkers directly implicated in MetS. These metabolites include glucose, triglycerides (also known as triacylglycerol or TAG), low-density lipoprotein cholesterol (LDL-C), and high-density lipoprotein cholesterol (HDL-C). We additionally chose to study the activity of the fatty acid beta-oxidation pathway. Although not a conventional marker, fatty acid oxidation plays a critical role in lipid metabolism, and the dysregulation of this pathway has been linked to MetS ^{12,13,14}.

We evaluated how different diets may influence the serum levels of glucose, TAG, LDL-C, and HDL-C by analyzing reactions in the WBMs involved in their transport (exchange) between organs and the bloodstream. For the male WBM, we extracted 29 exchange reactions for glucose, three for TAG, three for HDL-C, and three for LDL-C. For the female WBM, we extracted 30 exchange reactions for glucose, three for TAG, three for HDL-C, and five for LDL-C. For the fatty acid beta-oxidation pathway, we focused on the last step of this process, which involves the breakdown of butyryl-CoA into acetyl-CoA in mitochondria and extracted two reactions from both the male and female WBMs representing this conversion in key tissues contributing to this pathway. The complete list of these reactions is provided in **Supplementary Table 1.**

We performed separate in silico simulations for male and female using the sex-specific WBMs as typical male and female digital subjects. Computational simulations of the WBMs were performed using parsimonious FBA (pFBA) while constraining the whole-body maintenance reaction (see Methods).

Analysis of diets' nutrient composition and diversity

We analyzed the nutrient compositions of the 12 examined diets by employing Principal Component Analysis (PCA) using both their macro- and micronutrient compositions as features (**Figures 1B** and **1C**). Visualizing these diets using the first two principal components (PCs) revealed new insights into the relationships between these diets, shedding light on their underlying similarities and dissimilarities based on macro- and

micronutrient compositions. The macronutrient PCA (**Figure 1B**) highlights a spectrum of dietary profiles with the Balanced, DACH, High Fiber, and Mediterranean diets clustering towards a balanced macronutrient composition, indicative of a harmonious blend of carbohydrates, proteins, and fats. Conversely, the Keto diet with an extreme fat

Table 1 . Overview of the examined diets for this study.	
Diet	Description
Mediterranean Diet	The Mediterranean diet is made up of minimally processed foods, fresh fruits, and vegetables, and involves olive oil as its primary fat source. It assumes that dairy products are eaten daily, while poultry and fish are eaten occasionally, and red meat is eaten rarely.
Balanced Diet	The Balanced diet was designed to provide healthy amounts of essential nutrients, minerals, and vitamins to support normal metabolic functions for a healthy individual.
DACH Diet	The DACH diet, sourced from nutritional societies of Germany, Austria, and Switzerland. This diet was conceived to guarantee a nutritious health status for adults between the ages of 19 and 51 years old.
Vegetarian Diet	The Vegetarian diet is based on the most popular form of Vegetarianism, the ovo- lacto Vegetarian. This plant-based diet includes dairy and eggs but does not include any meat or fish.
Vegan Diet	The Vegan diet is another plant-based diet; however, it excludes all products derived from animals.
High Fiber Diet	The High Fiber diet contains animal sourced products and has a higher amount of dietary fiber than all other tested diets.
High Protein Diet	The High Protein diet is commonly used by athletes and those looking to lose weight.
Ketogenic (Keto) Diet	The Keto diet, also known as, the high fat low carb diet, aims to minimize the amount of carbohydrates intake and to replace them with fat and protein to promote the burning of body fat energy.
Gluten Free Diet	The Gluten Free diet excludes all foods containing gluten, a protein commonly found in wheat, rye, and barley.
Average European Diet	The components of this diet were based on an Austrian survey that was distributed in 2012 for approximately 1002 Austrian citizens across all age groups ⁴⁶ . This diet entails moderate intake of fat, protein and alcohol, and low intake of carbohydrates and fiber.
Average American Diet	The American diet comprises high levels of processed foods, red and processed meat, refined grains, simple sugars, saturated fats, and sodium.
Unhealthy Diet	The typical Unhealthy diet consists of high intake of kilocalories, simple sugars, saturated fatty acids, and cholesterol as well as a low amount of dietary fibers often sourced from fruits and vegetables.

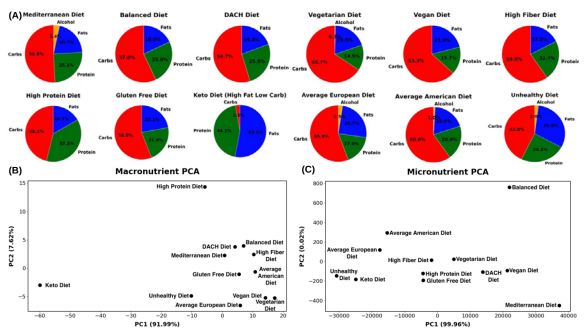


Figure 1. Macronutrient and micronutrient composition analysis of diets. (A) The macronutrient breakdown of the examined diets, (B) Spatial arrangement of diets based on their macronutrient, and (C) and micronutrient compositions. The diets were visualized using Principal Component Analysis (PCA), with reactions in the WBMs as features and diets as digital subjects (samples).

preference and very low carbohydrates uptake, shows an extreme deviation for the rest of the diets. Another outlier is the High Protein diet, which heavily emphasizes animal-derived protein sources and diverges notably from other diets. The Vegan and Vegetarian diets exhibited converging macronutrient compositions, emphasizing plant-based protein sources and a higher proportion of carbohydrates. The Average European diet is also positioned in proximity of the Vegetarian and Vegan diets, representing analogous proportions of carbohydrates and fats, coupled with a modest protein intake. The Average American and Gluten Free diets share similar profiles, suggesting comparable macronutrient distributions. Notably, the distinct positioning of the Average European diet in relation to its American counterpart reflects regional dietary preferences in Western diets. Finally, the Unhealthy diet's distinct position indicates a skewed macronutrient profile.

The micronutrient PCA plot presents a different perspective on these diets, diverging from that for macronutrients (**Figure 1C**). The Balanced and Mediterranean diets are both distinctly separated from each other and from the remaining diets, indicating their

unique micronutrient profiles. The High Fiber, Vegetarian, High Protein, DACH, Vegan, and Gluten Free diets aggregate into a pronounced cluster reflecting a congruence in their micronutrient compositions and potentially analogous impacts on metabolic processes and syndromes. The Unhealthy and Keto diets are clustered together on the left quadrant of the plot, representing their unique and similar micronutrient profiles that diverges from that of the healthier diets and may indicate potential nutrient deficiencies. In this landscape, the Average European diet is situated in the relative proximity of both the cluster of diverse diets (including the High Fiber and Vegetarian), and the cluster of Unhealthy and Keto diets, suggesting a partial overlap in micronutrient profiles with both groups. In contrast, the Average American diet remains notably separated and further from both clusters, implying a potentially distinct micronutrient profile that does not align with either of these diet clusters or with its European counterpart. Collectively, these analyses indicate the marked diversity in nutritional content across these diets.

The effect of diet on the systemic metabolic response

We examined how diverse dietary patterns influence the overall metabolic state in males and females. To this end, we utilized t-distributed Stochastic Neighbor Embedding (t-SNE) analysis with all the 81,094 and 83,521 metabolic reactions within the male and female WBMs, respectively. In these analyses, reactions served as features while the 12 diets served as samples with each diet corresponding to a digital subject on the respective diet. This analysis revealed the differential impact of these diets on the systemic metabolic response (Figure 2A and 2B). A discernible pattern is that the diets traditionally known to be healthier, including the Mediterranean, Balanced, Vegan, and DACH diets, manifest as a cluster in both males and females, reflecting their analogous and potentially beneficial modulation of metabolic state. Conversely, the Unhealthy and Keto diets are conspicuously isolated from these healthy diets and positioned on the opposing corners of the plots, implying a divergent influence on the systemic metabolism. The Vegetarian diet, while positioned separately, tends towards the healthier diets cluster in male and female, suggesting a favorable metabolic imprint akin to these diets. The High Protein, High Fiber, and Gluten Free diets occupy a middle locus in both males and females, which indicates a rather moderate influence on metabolic landscape, neither heavily favoring nor significantly diverging from a state of metabolic health. We also observe a notable dispersion of diets within these clusters, indicating a subtle gradation in their metabolic influence. Intriguingly, the Average American diet is clustered with the Average European diet and is located near the cluster of less healthy (Unbalanced and Keto) diets in males; however, it moves toward the cluster of healthier (Vegetarian, DACH, Balanced) diets in females.

The effect of diet on metabolic predisposition to MetS

To evaluate the interplay between diet and metabolic susceptibility toward developing MetS, we leveraged 35 and 43 reactions in the male and female WBMs, respectively, responsible for glucose, TAG, HDL-C, and LDL-C exchange with blood, as well as the reactions involved in fatty acid oxidation. These reactions were used as features for

visualization using t-SNE with the 12 diets serving as digital subjects (**Figures 2C** and **2D**). The t-SNE plot in males depicted a continuum of dietary impacts on metabolic processes pertinent to MetS with trends somewhat like those seen in **Figure 2A**, where healthier and unhealthy diets are clustered distinctly separated in the opposing corners of

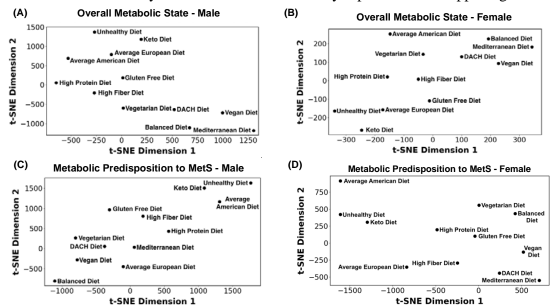


Figure 2. Visualization of the examined diets based on the systemic metabolic response and metabolic predisposition to MetS. (A) Systemic metabolic response for the male and (B) female WBMs. (C) Metabolic predisposition to MetS for the male and (D) female WBMs. The diets were visualized using t-Distributed Stochastic Neighbor Embedding (t-SNE), with reactions in the WBMs as features and diets as digital subjects (samples).

the t-SNE plot and the rest of the diets positioned between these two extremes. A noticeable divergence between the two is the repositioning of the Average European diet from the proximity of unhealthy diets in **Figure 2A** to the cluster of conventionally healthy diets in **Figure 2C**. This suggests a MetS risk profile for this diet similar to that of the healthy diets.

The t-SNE plot for females presents a rather distinct clustering of diets in relation to the MetS metabolic biomarkers compared to males (**Figure 2D**). A notable aggregation is evident in the lower right quadrant of the female t-SNE plot, comprising the Mediterranean, Vegan, and DACH diets. However, the other two healthy diets, namely the Balanced and Vegetarian diets, form an independent cluster in the upper right quadrant, representing a differentiated MetS risk profile compared to other healthy diets in females. While closer to the Unhealthy and Keto diets, the Average American diet is markedly set apart in the upper right corner, hinting at a unique MetS risk profile induced by this diet in females. Its European counterpart, however, along with the High-Fiber diet, is positioned closer to the cluster of healthier diets, similar to that in males. These

observations underscore the specific metabolic response of females to both healthy and unhealthy diets in relation to the MetS biomarkers.

The effect of diet on individual MetS biomarkers

In the subsequent sections, we evaluate the effect of diet on each molecular biomarker of MetS and fatty acid oxidation as well as the significance of different organs/tissues.

Glucose: Glucose is one of the most widely recognized diagnostic markers for MetS ¹⁵. Specifically, abnormally elevated fasting serum levels of glucose are considered a major MetS risk factor. We assessed a total of 24 reactions in males and 31 reactions in females involved in the transport of glucose between different organs/tissues and the systemic blood circulation.

Overall glucose secretion into the blood: Evaluating the total glucose secretion into the bloodstream (i.e., sum of glucose secretion fluxes by all organs secreting glucose) in the male WBM revealed that surprisingly the Balanced diet has the highest glucose secretion flux (3,994.7 mmol/person/day), while the Unhealthy diet (3,266.3 mmol/person/day) followed the Keto diet (3,728.9 mmol/person/day) and Average European diet (3,760.4 mmol/person/day) exhibit the lowest glucose section fluxes (Figure 3A). The rest of the diets displayed a consistent overall glucose secretion of 3,905.4 mmol/person/day in the male WBM. Similar patterns were observed in the female WBM, where, contrary to expectations, the healthier diets show the highest overall glucose secretion into the blood, while the unhealthy diet shows the reduced glucose secretion fluxes. Specifically, the Mediterranean diet shows the highest overall glucose secretion flux (7,228.0 mmol/person/day), followed by the Vegan (7,165.4 mmol/person/day), Balanced (7,161.0 mmol/person/day), and DACH (7,157.6 mmol/person/day) diets (**Figure 3B**). Conversely, the Unhealthy diet exhibits a significantly lower overall glucose secretion flux compared to all other diets (4,534.6 mmol/person/day). The Keto diet (5,371.4 mmol/person/day) and Average European (6,783.9 mmol/person/day) diets also revealed particularly low secretion levels similar to those in males.

Organs/tissues secreting glucose into the blood: Our analysis identified 18 organs/tissues in both male and female WBMs that consistently engaged in glucose secretion into the bloodstream (**Figure 3C**). The liver, thyroid gland, heart, adipocytes, and stomach emerged as major contributors to glucose secretion into blood in the male WBM. In the female WBM, the breast, uterus, and liver emerged as the organs with the highest contribution to glucose secretion into blood across all diets (**Figure 3D**).

Organs/tissues taking up glucose from the blood: Six organs/tissues in male and 11 in female WBMs demonstrated consistent glucose uptake from blood across all the diets. In males, the lungs emerged as the most avid consumer with a consistent glucose uptake flux of 2,417.4 mmol/day/person across all studied diets, followed by pancreas (1,000.3 mmol/day/person), and kidney (205.5 mmol/day/person) (**Figure 3C**). In females, the

lung, skin, and colon are responsible for taking up the highest levels of glucose from the blood across most diets (**Figure 3D**).

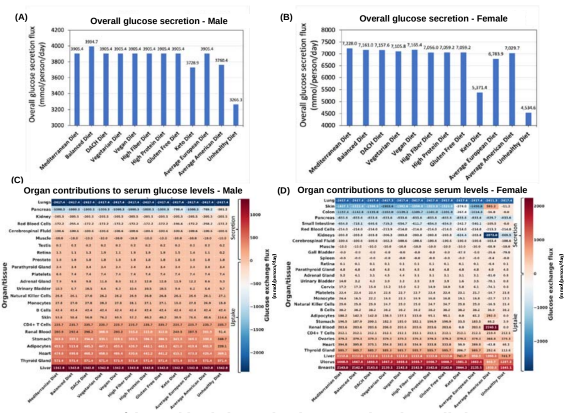


Figure 3. Impact of diet on blood glucose levels. (A) Predicted overall glucose secretion flux into the bloodstream for the male and (B) female WBMs. (C) Contribution of different organs/tissues within the male and (D) female WBMs to glucose secretion and uptake into/from the blood.

Organs with notable altered contributions to glucose serum levels across diets: The pancreas, adipocytes, and renal blood are among the organs/tissues in males for which their contributions to serum glucose levels are most influenced by diet variations (**Figure 3C**). The diets that induced the largest change in glucose secretion or uptake by these organs is the Unhealthy diet, followed by the Average American and Keto diets. Notably, these are the three diets that clustered together with respect to MetS risk (**Figure 2C**). For instance, the glucose uptake flux from the blood by the pancreas recorded to be 361.2, 769.2, and 799.4 mmol/day/person for the Unhealthy, Average American, and Keto diets, respectively, which are significantly lower than that for all other diets (1,000.3 mmol/day/person). Adipocytes, which act as the WBM's representation of fat storage distributed throughout the body, also experienced marked changes in glucose secretion in the male WBM under the Unhealthy, Average American, Balanced, and Vegan diets. The Balanced (513.0 mmol/day/person) and Vegan (455.6 mmol/day/person) diets both show

a notable increase in adipocytes glucose secretion flux into the blood relative to the other eight diets (433.0 \pm 16.0 mmol/day/person) (**Figure 3C**). Conversely, the Unhealthy diet followed by the Average American diet resulted in a substantial decrease in adipocytes' glucose secretion (220.1 and 401.9 mmol/day/person, respectively). The renal blood was another organ with altered glucose secretion profile across diets. In the WBMs, the kidney is associated with two distinct reactions involving the exchange of glucose with blood: the exchange of glucose between the kidney and systemic blood, and the exchange of glucose between the renal blood and systemic blood (depicted by rows 'Kidney' and 'Renal Blood', respectively in **Figure 3C**). Notably, the kidney is involved in glucose uptake from the systemic blood circulation, while the renal blood engages in glucose secretion into the systemic blood circulation. This can be explained by previous reports that the kidneys play a significant role in maintaining glucose homeostasis in the body by releasing glucose into the circulation via gluconeogenesis, taking up glucose from the circulation to fulfill their energy needs, and reabsorption of glucose at the proximal tubule ¹⁶. Here, we observed an increase in glucose secretion flux from renal blood into systemic blood under the High Protein and Gluten Free diets (both secretion fluxes were predicted at 312.0 mmol/day/person). In contrast, a significant decrease was observed under the Unhealthy diet (51.37 mmol/day/person) compared to the remaining nine diets $(300.9 \pm 18.5 \text{ mmol/day/person})$ (Figure 3C). No changes were observed under these diets for the glucose uptake from the systemic blood circulation by the kidney.

In females, the organs that are most affected by diet include the breast, uterus, liver, skin, and colon (**Figure 3D**). Specifically, the glucose secretion fluxes for the breast and uterus under the Average American and Unhealthy diets were significantly lower compared to other diets. Similarly, the liver exhibits a notable decrease in glucose secretion flux under the Keto, Unhealthy, Average European, and Average American diets compared to other diets—a pattern not observed in the male WBM. Skin and colon also experience diminished glucose uptake from the blood under the Unhealthy, Average American, Keto, and Average European diets. Notably, we noticed several outlier responses for glucose exchange between blood and organs under the Average American diet for the female WBM. For example, while skin cells engage in glucose uptake from blood under all other diets, they show glucose secretion rather than uptake under the Average American diet. Similar abnormalities were observed in the retina, adrenal gland, urinary bladder, cervix, platelets, monocytes, NK cells, and the heart.

TAG: Formed by the esterification of three fatty acid molecules to glycerol, TAG is a prevalent form of fat and primary energy storage within the human body. Elevated serum levels of TAG are a hallmark of MetS ^{15, 17}. Elevated serum levels of TAG are often associated with the consumption of a high-calorie diet rich in refined carbohydrates and saturated fats, which can contribute to dyslipidemia— another leading factor implicated in MetS ⁴. TAG biosynthesis occurs predominantly in the liver, where excess dietary carbohydrates and proteins—surpassing the body's immediate needs—are converted into fatty acids and subsequently into TAG. Concurrently, the small intestine is instrumental in the absorption of dietary fats, predominantly in the form of TAG. Before entering the

bloodstream, TAG is packaged into very low-density lipoproteins (VLDL) in the liver and chylomicrons in the intestine as it is not soluble in the aqueous environment of the blood. Adipose tissues serve as the principal depots for TAG storage. Upon arrival via the bloodstream, TAG is hydrolyzed back into free fatty acids and glycerol, mediated by the

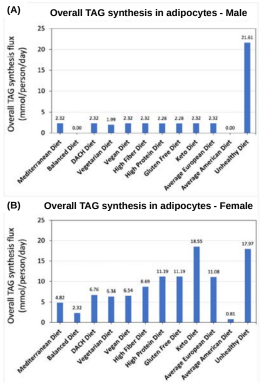


Figure 4. Impact of diet on TAG synthesis. (A) Predicted overall TAG synthesis in the adipocytes in the male and (B) female WBMs.

enzyme lipase. The liberated fatty acids and glycerol are then absorbed by adipocytes and re-esterified into TAG for long-term storage ¹⁷.

In our analyses, we assessed TAG storage within adipocytes in the WBMs as a proxy for the TAG serum levels. This was accomplished by examining the flux of three specific TAG synthesis reactions in adipocytes within both the male and female WBMs, each responsible for the esterification of different types of essential and non-essential fatty acids into TAG within the adipocytes. We computed the aggregate flux through these three reactions, treating this sum as a representation of total TAG synthesis and accumulation within adipocytes.

Our analysis revealed a pronounced disparity in TAG synthesis and storage within adipocytes across various diets in both male and female WBMs. In males, the Unhealthy diet exhibits a considerably higher TAG synthesis flux of 21.61 mmol/day/person compared to the remaining diets (1.858 \pm 0.923 mmol/day/person) (**Figure 4A**), which

can be attributed to the diet's excessive calorie load and unhealthy fats intake. Conversely, the Balanced diet is characterized by a negligible TAG synthesis, suggesting a dietary composition that mitigates excessive lipid accumulation. Contrary to expectation, the Average American Diet, despite its reputation for high caloric content and associations with MetS, also shows a negligible TAG synthesis in adipocytes. The rest of the diets show comparable levels of TAG synthesis flux in adipocytes although Vegetarian (1.99 mmol/day/person), High Protein (2.28 mmol/day/person), and Gluten Free diets (2.28 mmol/day/person) exhibit slightly lower TAG synthesis and storage flux relative to the remaining diets.

Our analysis using the female WBM revealed that TAG synthesis and storage in adipocytes was highest under the Keto (18.55 mmol/person/day) and Unhealthy diets (17.97 mmol/person/day) (**Figure 4B**). The high fat content of the Keto diet (more than 53%; **Figure 1**) appears to significantly increase the TAG synthesis in females, a pattern diverging from that for the males. Again, consistent with males but counterintuitively, the Average American shows the lowest TAG synthesis in females (0.81 mmol/person/day), The next diets with low TAG synthesis and storage in adipocytes are the Balanced (2.32 mmol/person/day), Mediterranean (4.82 mmol/person/day), and Vegetarian (6.34 mmol/person/day) diets, aligning with our expectation for these healthier diets.

LDL-C and **HDL-C**: HDL-C and LDL-C are two types of lipoproteins that transport cholesterol throughout the body via blood circulation thereby regulating cholesterol levels. LDL-C (often referred to as "bad cholesterol") is responsible for transporting cholesterol from the liver to various tissues within the body ¹⁸. HDL-C (also known as the "good cholesterol") engages in reverse cholesterol transport, removing excess cholesterol from the bloodstream and peripheral tissues and transporting it back to the liver for disposal. Elevated levels of LDL-C can lead to the build-up of cholesterol in the arteries, forming plaques that can narrow down and obstruct blood vessels—a condition referred to as atherosclerosis, which increases the risk of MetS and cardiovascular disease. Conversely, high serum levels of HDL-C, have a protective effect against atherosclerosis¹⁹, while its low serum levels contribute to the increased prevalence of MetS, particularly in males ²⁰.

In our analyses, we focused on three distinct reactions for HDL-C and three others for LDL-C within the male WBM that contribute non-zero (although sometimes very small) flux to HDL-C and LDL-C exchange with blood within the WBMs—namely the liver, muscle, and adipocytes. For the female WBM, we selected three reactions for HDL-C and five reactions for LDL-C that exchange HDL-C and LDL-C between the bloodstream and liver, muscle, adipocytes, kidney, and renal blood.

Overall LDL-C/HDL-C blood secretion ratio: We first evaluated the total secretion flux of HDL-C and LDL-C by all relevant organs for each diet and then calculated the ratio of total LDL-C to HDL-C secretion fluxes (LDL-C/HDL-C). For the male WBM, this analysis revealed that the Unhealthy diet exhibits a substantially higher LDL-C/HDL-C ratio (1.54) compared to all other diets (**Figure 5A**), indicating impaired cholesterol

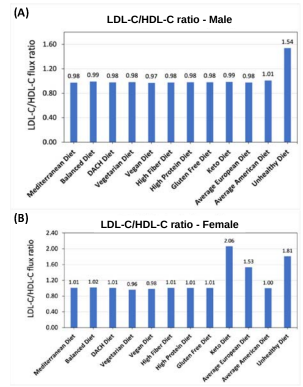


Figure 5. Effect of diet on LDL-C to HDL-C ratio. (A) Predicted LDL-C/HDL-C ratio in the male and (B) female WBMs.

metabolism and potentially increased risk for MetS. The ratios for the rest of the diets are comparable although there are slight variations (0.98 \pm 0.01). The Average American diet shows the second highest LDL-C/HDL-C ratio (1.01), which underscores the diet's association with adverse lipid profiles and heightened MetS risk. In contrast, the Vegan diet shows the lowest LDL-C/HDL-C ratio (0.97) in males, indicating a more favorable lipoprotein profile. The remaining diets exhibit intermediate LDL-C/HDL-C ratios, with the DACH, Vegetarian, High Fiber, High Protein, Gluten Free, and Keto diets having a consistent LDL-C/HDL-C ratio of 0.98 and the Balanced and Mediterranean having a ratio of 0.99. This reflects a moderate influence of these diets on cholesterol metabolism.

When examining the LDL-C/HDL-C ratio in females, we found the greatest ratios under the Keto (2.06) and Unhealthy (1.81), and Average European (1.53) diets. Conversely, the lowest LDL-C/HDL-C ratios occurred under the Vegetarian (0.96) and Vegan (0.98)

diets (**Figure 6B**). The remaining diets (Mediterranean, Balanced, DACH, High Fiber, High Protein, Average American, and Gluten Free diets) revealed mostly intermediate ratios ranging from 1.00 to 1.02. Again, we expected a higher LDL-C/HDL-C ratio for the Average American diet, considering its high intake of saturated fats.

Fatty acid beta-oxidation pathway: Fatty acid beta-oxidation pathway plays a pivotal role in the human body's energy metabolism by breaking down fatty acids to generate ATP. This pathway's efficiency is crucial for maintaining metabolic health as deficiencies can lead to elevated free fatty acid levels, which contributes to metabolic diseases including MetS ^{21, 22}. However, the complexity of this metabolic process and its limited experimental accessibility in human cells has hampered a deep understanding of the pathophysiological mechanisms underlying the link between dietary factors, dysregulation of this pathway, and the risk of metabolic diseases, especially MetS ^{21, 22}.

We explored the effect of various diets on the activity of fatty acid beta-oxidation pathway. Our analysis centered on the final step of this metabolic process, the conversion of butyryl-CoA, a 4-carbon fatty acid, into acetyl-CoA, a 2-carbon molecule, catalyzed by the enzyme thiolase. The flux of this reaction was used as a surrogate for the activity of the fatty acid oxidation process. Although fatty acid beta-oxidation occurs in all tissues and organs within the human body, the (skeletal) muscles and heart (cardiac muscles) are the major contributors ²³. Therefore, we focused our investigations especially on the muscles and heart in both the male and female WBMs. This rationale guided us to select two specific reactions within the male and female WBM representing butyryl-CoA breakdown in these tissues/organs.

Overall fatty acid beta-oxidation activity: We first evaluated the overall fatty acid beta-oxidation activity within the human body as captured by sum of the butyryl-CoA breakdown fluxes in the muscles and heart. For the male WBM, the Vegan (67.2 mmol/person/day) and Vegetarian (67.0 mmol/person/day) diets followed by the Average European (64.5 mmol/person/day) and Keto (64.1 mmol/person/day) diets exhibited the highest overall activity (**Figure 6A**). On the other end of the spectrum, unexpectedly the Balanced diet followed by the Average American diets showed the lowest activity (44.0 and 49.4 mmol/person/day, respectively). The remaining diets show intermediate activity.

For the female WBM, we observed a substantially higher activity of fatty acid beta-oxidation for the Average American diet (76.0 mmol/person/day) compared to other diets (**Figure 6B**)—a pattern contrary to what we observed for the male WBM. The Balanced (40.8 mmol/person/day Vegan (31.4 mmol/person/day), Mediterranean (31.0 mmol/person/day), and DACH (28.5 mmol/person/day) diets are the next diets high fatty acid oxidation activity. Conversely, the Unhealthy and, unexpectedly, the Keto (7.4 mmol/person/day, respectively) diets demonstrated the lowest activity.

The effect of diets on organs' contributions to fatty acid oxidation: In the male WBM, the highest fatty acid oxidation activity in muscles was observed under the Vegetarian diet (35.7 mmol/person/day) and the Keto (34.8 mmol/person/day) diets (**Figure 6C**). The lowest activity in muscle was observed for the Balanced (19.6 mmol/person/day) followed by the Average American (24.0 mmol/person/day) and Unhealthy (28.9. mmol/person/day) diets. Similar observations were made for the heart which, like muscle,

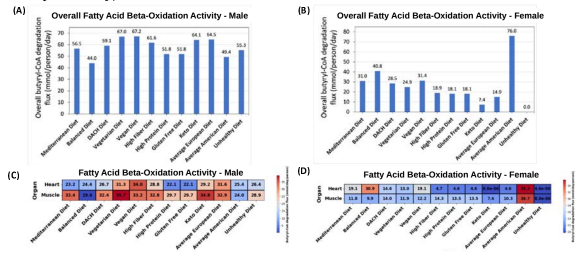


Figure 6. Effect of diet on the fatty acid beta-oxidation pathway. (A) Predicted overall fatty acid oxidation activity for the male and (B) female WBMs. (C) Contribution of the muscles and heart to fatty acid oxidation in the male and (D) female WBMs.

depends on fatty acids as a primary energy source—a process is crucial for the heart's constant contraction and efficient functioning ²⁴. For the heat the Vegan (34.0 mmol/person/day) diet followed by the Average European and Vegetarian (31.6 and 31.3 mmol/person/day, respectively) diets exhibited the highest activity. Conversely, the High Protein and Gluten Free diets (22.1 mmol/person/day for both), along with the Mediterranean (23.2 mmol/person/day) and Balanced (24.4 mmol/person/day) diets exhibited the lowest activities.

For the female WBM, the Average American diet induces a substantially higher fatty acid oxidation activity in both the muscles (36.7 mmol/person/day) and heart (39.3 mmol/person/day) compared to other diets (**Figures 6C** and **6D**). The High Fiber (14.3 mmol/person/day) and DACH (14.0 mmol/person/day) diets exhibit the next highest activity in the muscles. This is supported by existing literature that dietary fiber can induce metabolic adaptations in the muscles, potentially upregulating the activity of fatty acid oxidation ^{25,26}. For the heart, the Balanced, Mediterranean, and Vegan diets lead to the next highest fatty acid oxidation activity (30.9, 19.1, and 19.1 mmol/person/day, respectively). Conversely, the lowest activity in both the muscles and heart was observed for the Unhealthy and Keto diets.

Discussion

We systematically evaluated the impact of 12 diverse dietary regimens on key biomarkers associated with MetS through the lens of sophisticated organ-resolved whole-body models of metabolism. Our investigation explored the secretion of glucose, TAG, LDL-C, and HDL-C into the bloodstream, as well as fatty acid oxidation activity in both male and female. This in silico analysis revealed how specific dietary patterns influence the physiological processes underlying these MetS biomarkers and uncovered new insights into the contribution of different organs and tissues.

Our study supports the existing hypotheses that unhealthy diets—characterized by low in dietary fibers and high in saturated fats, cholesterol, and simple sugars—tend to adversely impact MetS biomarkers, while healthy diets, i.e., those involving the intake of whole grains, fruits, vegetables, lean proteins, and healthy fats promote metabolic homeostasis and reduce MetS risk. Nonetheless, we also identified counterintuitive relationships between specific diets and MetS biomarkers, indicating the complex interplay between dietary intake and physiological processes underlying MetS.

The analysis of diets according to nutrient composition illustrates high variability among diets and highlights the importance of considering both macro and micronutrient profiles: PCA provided a macroscopic view of the dietary patterns examined in our study, revealing distinct clusters of diets based on their macronutrient (Figure 1B) and micronutrient compositions (Figure 1C). This analysis illustrated a gradient of macro and micronutrient diversity and density in the examined diets, with distinct spatial distribution of diets in the micronutrient PCA plot compared to macronutrients. Specifically, despite both being considered healthy diets, the Mediterranean and Balanced diets occupy conspicuously isolated positions in the micronutrient PCA plot, separate from each other and from the remaining diets—a pattern not observed in the macronutrient PCA plot. This suggests a unique micronutrient profile for these diets, diverging from the remaining diets. The Mediterranean diet, rich in plant-based foods, olive oil, and fish, likely provides a distinct array of vitamins, minerals, and phytonutrients. The Balanced diet's unique positioning may also reflect its diverse and equilibrated nutrient composition that adheres to nutritional guidelines for promoting metabolic health. The Average American diet also stands apart from the cluster of unhealthy diets in the micronutrient PCA plot, emphasizing its unique micronutrient composition. These observations highlight the importance of considering both macro and micronutrient compositions when investigating the effect of dietary regimens on metabolic health.

Examining the impact of diet on systemic metabolic responses and overall MetS predisposition, revealed clear separations between healthy and unhealthy dietary patterns across both genders: Our t-SNE analysis offered a deeper dive into the effects of these diets on both systemic metabolic response (Figures 2A and 2B) and metabolic predisposition to MetS (Figures 2C and 2D) in male and female. Analysis of metabolic response at the systemic level revealed that diets traditionally considered healthy—such

as the Mediterranean, Vegan, DACH, and Balanced—formed clusters that significantly diverged from those deemed less healthy, like the Unhealthy and Keto diets, in both male and female. A similar divergence of the healthy and unhealthy diet clusters was also observed in the t-SNE plots representing metabolic susceptibility to MetS. This spatial stratification highlights the notably distinct impacts of these dietary groups on both the systematic metabolic homeostasis and MetS. It also demonstrates the marked potential of diet to modulate the biochemical milieu associated MetS. The consistent clustering of the Keto diet with the Unhealthy diet suggests potential adverse metabolic outcomes despite its popularity for weight loss. In addition to these commonalities between the systematic metabolic response and MetS susceptibility, we noticed differences between the two as well. A notable deviation between the two is the repositioning of the Average European diet from the close proximity of the Unhealthy diet when examining systematic metabolic response (Figures 2A and 2B) to that of the cluster of healthier diets when exploring response to MetS risk (Figures 2C and 2D) in both the male and female WBMs. This indicates a MetS risk profile of this diet that is closer to the healthier diets, in contrast to its American counterpart showing close proximity with respect to both systematic metabolic response and MetS risk.

Unhealthy diets yield paradoxically lower glucose secretion profiles despite higher glycemic loads compared to healthier diets: Analyzing the impacts of diet on individual MetS biomarkers using the WBMs provided granular insights into the complex interactions between dietary intake and the physiological indicators. For instance, when examining the effect of diet on glucose secretion into the systemic blood, the Unhealthy, Keto, and Average American, diets resulted in lower overall glucose secretion into the blood compared to healthier diets in both the male and female WBMs (Figures 3A and **3B**). Specifically, the Balanced and the Unhealthy diets showed the highest and lowest glucose secretion, respectively, in males. Similar patterns were observed in females, where the Mediterranean diet led to the highest glucose secretion and the Unhealthy diet showed the lowest. These results contrast with our expectations that unhealthy diets are presumed to elevate glucose levels due to higher glycemic loads. This counterintuitive finding might be attributed to the body's complex metabolic processes and its ability to maintain homeostasis. For instance, the body might be compensating for the high intake of simple sugars in unhealthy diets by reducing glucose secretion. Additionally, the composition of these unhealthy diets, often high in processed foods and simple sugars, could cause a rapid spike in blood glucose levels followed by a sharp decline, leading to lower overall glucose secretion. These findings could be also influenced by the inherent limitations of the WBMs. Importantly, these observations suggest that the risk of MetS may not solely be determined by the healthiness of a diet but also by how the body metabolically responds to different dietary patterns. Further experimental studies are needed to validate these computational findings.

Organ-specific contributions to blood glucose homeostasis reveal the critical role of the liver and lungs, highlighting their pivotal influence on MetS risk: The organs identified to play a major role in blood glucose levels align with established physiological

knowledge. For instance, the liver, a known site for gluconeogenesis (glucose production from non-carbohydrate precursors) and glycogenolysis (glycogen breakdown to glucose), emerged as a major glucose secreting organ in both males and females (**Figures 3C** and **3D**). Our study also identified the lungs as a major contributor to glucose blood levels, exhibiting the highest glucose uptake from the blood in both males and females, although the uptake levels remained unaffected by dietary variations. This major role of the lungs in glucose metabolism is consistent with previous reports ²⁷ and implies that any impairments in the lung function can critically affect serum glucose levels and, therefore, raise the risk of developing MetS. This finding is supported by existing evidence that lung conditions such as asthma and pulmonary hypertension are associated with abnormally high postprandial blood glucose levels ²⁸. In addition, Baffi et al. ²⁹. emphasize that MetS is closely linked to lung health. These highlight that the lung's role in managing the blood glucose homeostasis and MetS risk is crucial yet often overlooked.

Dietary variations, especially diets high in fats and simple sugars, markedly influence blood glucose contributions from renal blood, pancreas, and adipocytes in males, and from breasts, uterus, and liver in females: Interestingly, our findings revealed that specific organs are more responsive to dietary changes than others. In males, the renal blood, pancreas, and adipocytes exhibited the most significant response to dietary variations. In particular, we observed an increase in glucose secretion from renal blood into the systemic blood under the High Protein and Gluten Free diets and a notable decrease under the Unhealthy diet. While the reason behind this dramatic decrease in glucose secretion under the Unhealthy diet has not been reported before, the observed increase for the High Protein diet has been documented. For example, protein feeding in mice has been reported to induce an increase in endogenous glucose production in the kidney and consequently increased glucose release into blood compared to a normal starch diet ³⁰. Another striking finding was the significant reduction in pancreatic glucose uptake under the Unhealthy, Average American, and Keto diets compared to other diets in males (Figure 3C). The pancreas plays a critical role in blood sugar regulation by secreting insulin. Increased glucose uptake by pancreatic cells is necessary for proper insulin production and secretion. Our observations thus suggest a potential link between unhealthy diets high in saturated fat, simple sugars, and processed foods and impaired pancreatic metabolic function. In females, notable variations in organ contributions to blood glucose levels were observed in the breast and uterus for the Average American and Unhealthy diets, with significant reductions in glucose secretion compared to other diets. The liver also showed decreased glucose secretion under the Keto and Unhealthy diets while the skin and colon experienced diminished glucose uptake from the blood under the Unhealthy, Keto, Average American, and Average European diets—further emphasizing the gender-specific metabolic responses to diet.

Pronounced differences in TAG synthesis across diets reflect the influence of high-caloric and fat as well as high-fiber and balanced content: Our analysis of TAG synthesis in adipocytes as surrogate for TAG serum levels revealed pronounced differences across diets (**Figure 4**). In males, the Unhealthy diet exhibited a substantial

increase in TAG synthesis compared to other diets, reflecting its high caloric content and unhealthy fats, which contribute to lipid accumulation. This finding aligns with previous studies showing that diets rich in saturated fats and refined sugars promote adipogenesis and lipid storage ^{31,32}. Conversely, the Balanced diet exhibited negligible TAG synthesis, highlighting its role in preventing excessive lipid accumulation. Vegetarian, High Protein, and Gluten Free diets also show slightly reduced TAG when compared to other diets. This aligns with existing literature suggesting that high protein and complex carbohydrate intake —a characteristic of these three diets— promote satiety, reduce calorie intake, and may improve glycemic control by reducing blood sugar spikes after meals, all of which could contribute to decreased TAG synthesis ³³. In females, the Keto and Unhealthy diets led to the highest while the Balanced, Mediterranean, and Vegetarian diets showed the lowest TAG synthesis fluxes. These findings support the adverse impact of high-caloric and high-fat diets and the key role of balanced, fiber-rich diets in regulating TAG storage in adipocytes.

Differential LDL-C/HDL-C ratios across diets highlight the adverse effects of highcalorie and high-fat diets and the favorable impacts of plant-based diets on lipid **profiles**: When examining cholesterol metabolism, the analysis of LDL-C/HDL-C ratio provided critical insights. The Unhealthy diet exhibited the highest LDL-C/HDL-C ratio in males (Figure 5A), indicating impaired cholesterol metabolism and increased MetS risk. This finding is supported by evidence linking high saturated fat intake with adverse lipid profiles ^{34,35,36}). Conversely, the Vegan diet had the lowest LDL-C/HDL-C ratio, reflecting a lower MetS risk. In females, the Keto diet followed by the Unhealthy diet also resulted in the highest LDL-C/HDL-C ratios. The high ratio for the Keto diet suggests that this diet adversely affects lipid profiles more severely in females compared to males. Similar to males, the Vegetarian and Vegan diets exhibited the lowest ratios in females (**Figure 5B**) These diets are typically lower in saturated fats and higher in dietary fibers and unsaturated fats, which promote the synthesis of HDL-C and the clearance of LDL-C from the bloodstream. The low LDL-C to HDL-C ratios for these diets support the potential cardioprotective effects associated with plant-based dietary patterns and their role in reducing the risk factors for MetS ³⁷.

Dietary influence on fatty acid beta-oxidation demonstrates the potential of plant-based diets to rival high-fat regimens in enhancing lipid metabolism: In addition to the four conventional biomarkers implicated in MetS, we also studied the activity of the fatty acid beta-oxidation pathway by focusing on the conversion of butyryl-CoA to acetyl-CoA, the last step in this pathway as a proxy. While not a conventional marker for MetS, this pathway plays a significant role in lipid metabolism, the dysregulation of which is a known factor in the development of MetS ²². In males, the Vegan and Vegetarian diets exhibited the highest fatty acid oxidation activity (**Figure 6A**), likely due to the increased intake of medium-chain fatty acids (MCFAs), which are prevalent in these diets and have been associated with increased fatty acid oxidation ³⁸. The Average European Diet followed by the Keto diet also showed high activity. This observation for the Keto diet was expected due to its high-fat and low carbohydrate content, which shifts

energy metabolism towards increased lipolysis and fatty acid utilization. This leads to the production of ketone bodies that provide an alternative energy source for the body when carbohydrate intake is low ³⁹. Our findings also suggest that promoting lipolytic activity within the body is not exclusively dependent on the adoption of extreme dietary regimens, such as the Keto diet; rather, a plant-centric diet may offer comparable, or even superior, efficacy. On the other end of the spectrum, the Balanced and Average American diets showed the lowest activity, suggesting a metabolic shift towards other energy sources. In females, we observed divergent patterns where paradoxically the Average American diet, followed by the Balanced and Vegan diets had the highest activity, while the Unhealthy and unexpectedly Keto diets exhibited the lowest activity (**Figure 6B**). These results reinforce the effectiveness of plant-based diets in promoting lipid oxidation.

Gender differences in metabolic responses to diets highlight the need for tailored nutritional strategies: Throughout our analysis, we observed several commonalities in the metabolic responses to dietary changes in both males and females as described above, suggesting that these responses are largely independent of gender. However, a number of distinct gender-specific patterns also emerged. For instance, when examining the systemic metabolic response to diet, we noticed that Average American diet clusters with the Average European diet and is proximal to less healthy diets such as the Unbalanced and Keto diets in males (Figure 2A); however, it shifts towards the cluster of healthier diets—Vegetarian, DACH, Balanced—in females (Figure 2B). Furthermore, the female WBM reveals a rather distinct clustering of healthier diets in relation to MetS biomarkers compared to males (Figures 2C and 2D). Specifically, while some healthy diets like the Mediterranean, Vegan, and DACH aggregate in the lower right quadrant of the t-SNE plot, others (the Balanced and Vegetarian diets) form a separate cluster in the upper right quadrant, indicating a differentiated MetS risk profile—a trend not observed in males.

In regard to lipid metabolism, the Average American diet shows the second highest LDL-C/HDL-C ratio in the male WBM, but it exhibits the third lowest ratio in the female WBM (**Figure 5**). This is consistent with prior studies reporting that while high and low levels of LDL-C influence the prevalence of MetS in males, females are affected at a lesser extent from high serum levels of LDL-C and are not affected by low serum levels ²⁰. Regarding fatty acid oxidation, the Balanced and Average American diets are associated with the lowest activity in males (**Figure 6A**), while in females, a reversed pattern emerges where these diets exhibit the highest oxidation activity (**Figure 6B**).

Notably, the Keto diet demonstrates divergent trends between genders with regard to the MetS biomarkers. For instance, it induces substantially higher TAG synthesis in females (Figure 4B)—a pattern not observed in males. This finding is corroborated by existing literature that high levels of TAG may be more strongly associated with the increased risk of MetS in females compared to males ⁴⁰. Moreover, while the Keto diet promotes high fatty acid oxidation in males (Figure 6A), it shows the second lowest activity in females (Figure 6B). This suggests that the Keto diet may not be an effective strategy for weight loss and lowering the risk of MetS in females.

The Average American diet also presented several counterintuitive results, especially for the female WBM. Notably, unexpectedly low TAG synthesis and storage were observed in both male and female models under this diet (Figures 4). Additionally, this diet displayed the highest fatty acid oxidation activity in the female WBM (Figure 5B). This unexpected outcome may reflect the diet's high content of saturated fats and refined carbohydrates, which, despite their association with negative health outcomes, could inadvertently fuel an increase in fatty acid metabolism in females. The low TAG synthesis along with high fatty acid oxidation activity might also indicate a compensatory metabolic mechanism by the body aimed at managing the excessive intake of saturated fats in this diet. Other unexpected metabolic responses to the Average American diet in females, further complicating our understanding of its metabolic impact include one of the lowest LDL/HDL ratios (Figure 6B), and abnormal glucose exchange patterns in female organs such as the skin, retina, and adrenal gland (Figure 2D). These findings may reflect an unconventional metabolic response and metabolic dysregulation in females that is unique to this diet's specific nutrient profile. This hypothesis is supported by the unique positioning of the Average American diet in females t-SNE plot (Figure **2D**). These findings may also hint at potential limitations of the female WBM.

Collectively, these observations indicate the sex-specific metabolic processing of the micronutrient content in diets and consequently specific metabolic response of males and females to both healthy and unhealthy diets in relation to the MetS risk. Thes also underscores the need for gender-specific dietary recommendations to mitigate MetS risk.

Limitations of this study: It is important to also highlight some important limitations of the present study. While the WBMs utilized in this investigation incorporate all the major organs/tissues involved in the metabolic processes relevant to MetS, the resolution is limited to organ-level processes, as these models were constructed using bulk RNA sequencing data from these organs/tissues. These models, therefore, do not account for the heterogeneity and specific functions of many distinct cellular types within each organ, potentially oversimplifying complex biological interactions. We also lacked the ability to capture metabolite concentrations in the blood as the WBMs are stoichiometric models that can predict reaction fluxes only. Consequently, changes in serum concentrations of target metabolites in response to dietary variations were not directly observable.

Another critical constraint of our studies is the exclusion of broader physiological regulatory mechanisms such as hormone activity, allosteric regulation of enzymes, signal transduction, and gene regulation, which are not captured within the WBMs. This omission might lead to an incomplete portrayal of the diet's influence on MetS biomarkers. For instance, most people with MetS also suffer from insulin resistance ⁴¹ making it more difficult for tissues/organs in the body to respond to insulin and to uptake glucose, but these effects cannot be captured within the WBMs.

These limitations necessitate a cautious interpretation of our findings as they are confined to the specific scope of the male and female WBMs, which focus solely on metabolic pathways and reaction fluxes and exclude non-metabolic regulatory processes that significantly influence physiological responses to dietary inputs. Therefore, further experimental validation in future studies is necessary to corroborate some findings from our studies that are not supported by the existing literature.

Some predictions by the WBMs might be interpreted in light of these limitations. For example, while both the male and female WBMs predict glucose, LDL-C, HDL-C secretion by adipocytes under the examined diets, current literature does not support direct glucose, LDL-C, and HDL-C secretion by adipocytes into the bloodstream, Adipocytes are, however, recognized to play a pivotal role in maintaining glucose homeostasis and regulating insulin sensitivity ^{42,43}. Additionally, they play a key role in the lipid metabolism and cholesterol efflux ⁴⁴, thereby influencing the blood levels of these lipoproteins. The inclusion of glucose, LDL-C, and HDL-C exchange reactions between the adipocytes and blood in WBMs, along with their predicted secretion into the blood could thus reflect an oversimplification of the adipocytes' role in glucose and lipid metabolism in these models.

Conclusion

In this study, we presented a rigorous in silico investigation that contributes to a deeper understanding of the impact of various dietary regimens on the risk of developing MetS. Our findings not only revealed novel insights into the molecular mechanisms underlying the relationships between poor diets and elevated risk of MetS, but also challenge some established notions about diet and metabolic health. Notably, this study underscores the importance of considering the micronutrient composition of diets and gender differences in response to dietary interventions. This aligns with the emerging trend in nutritional science advocating for personalized dietary interventions based on individual metabolic profiles ⁴⁵. Although further empirical research is needed to confirm our findings, our study demonstrates the potential of in silico modeling in elucidating intricate biological responses to dietary interventions, which are challenging to parse using in vitro or in vivo studies. For instance, by using this approach we were able to computationally identify the contribution of each organ/tissue toward the metabolic processes necessary for metabolic homeostasis or disruption thereof in MetS. Furthermore, the use of in silico models can mitigate the influence of confounding environmental variables typically seen in dietary intervention studies, thereby offering clearer insights into the metabolic reprogramming associated with MetS. Overall, our findings have the potential to lead to a more comprehensive understanding of metabolic health and diet and to inform more effective dietary intervention strategies to manage or mitigate the risk of MetS.

Materials and Methods

Whole-body models: We utilized sex-specific WBMs constructed by Thiele et. al ¹⁰ and constrained with metabolomic and physiological. The latest versions of the male and

female WBMs were obtained from the Virtual Metabolic Human (VMH) database ⁴⁶. The male model captured a total of 81,094 reactions and 56,452 metabolites, representing a typical male subject with a body weight of 70 kg, a height of 170 cm, a consistent heart rate of 67 beats per minute, a stroke volume of 80 mL, a cardiac output of 5360, and hematocrit of 0.400 ⁴⁶. The female model captured a total of 83,521 reactions and 58,851 metabolites, a body weight of 58 kg, height of 160 cm, and with all remaining parameters having the same values as the male model ⁴⁶.

In silico diets: The in silico diet formulations for the Average European, DACH, High Protein, High Fiber, Mediterranean, Unhealthy, Keto, Gluten Free, Vegetarian, and Vegan diets were obtained from the pre-defined diets section on the VMH database ⁴⁶. The Average American diet is based on the average diet of American males ages 20 and older according to a 2007-2008 NHANES Study and its in silico formulation was obtained from a previous study ⁴⁷. The in silico composition of the Balanced diet was also adopted from Sahoo and Thiele ⁴⁸.

Computational simulations of WBMs: We performed the computational simulation of the WBMs under different diets by using parsimonious Flux Balance Analysis (pFBA) where the Euclidean norm of reaction fluxes in the model was minimized as the objective function, following that in Thiele et. al ¹⁰. Other than the bounds on uptake reactions for dietary compounds, the rest of the lower and upper bounds on internal reactions and all the constraints were the same as those in the WBMs ¹⁰. Computational simulations were conducted in MATLAB 2019b (MathWorks Inc) by utilizing the COBRA (Constraint-Based Reconstruction and Analysis) Toolbox ⁴⁹. We used IBM ILOG CPLEX Optimization Studio V12.10.0, to solve the pFBA optimization problem. All simulations were performed on a local computer with a 3.3 GHx Dual-Core Intel i7 processor and 16 GB of memory.

Dimension reduction techniques: We conducted dimension reduction analyses using MATLAB 2019b's PCA and t-SNE functions (with default parameters for t-SNE), which are part of the Statistics and Machine Learning toolbox for MATLAB.

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Author contributions

ARZ conceived the study and, together with DSA, interpreted the results and drafted the manuscript. DSA performed all analyses. CVM assisted with computational simulations and provided the in silico Average American diet formulation. All authors have read and approved the final manuscript.

References

- 1. Rochlani Y, Pothineni NV, Kovelamudi S, Mehta JL. Metabolic syndrome: pathophysiology, management, and modulation by natural compounds. Ther Adv Cardiovasc Dis. 2017;11(8):215-25. Epub 20170622. doi:
- 10.1177/1753944717711379. PubMed PMID: 28639538; PMCID: PMC5933580.
- 2. Saklayen MG. The Global Epidemic of the Metabolic Syndrome. Curr Hypertens Rep. 2018;20(2):12. Epub 20180226. doi: 10.1007/s11906-018-0812-z. PubMed PMID: 29480368; PMCID: PMC5866840.
- 3. Noubiap JJ, Nansseu JR, Lontchi-Yimagou E, Nkeck JR, Nyaga UF, Ngouo AT, Tounouga DN, Tianyi FL, Foka AJ, Ndoadoumgue AL, Bigna JJ. Geographic distribution of metabolic syndrome and its components in the general adult population: A meta-analysis of global data from 28 million individuals. Diabetes Res Clin Pract. 2022;188:109924. Epub 20220515. doi: 10.1016/j.diabres.2022.109924. PubMed PMID: 35584716.
- 4. Castro-Barquero S, Ruiz-León AM, Sierra-Pérez M, Estruch R, Casas R. Dietary Strategies for Metabolic Syndrome: A Comprehensive Review. Nutrients. 2020;12(10). Epub 20200929. doi: 10.3390/nu12102983. PubMed PMID: 33003472; PMCID: PMC7600579.
- 5. Rozendaal YJW, Wang Y, Paalvast Y, Tambyrajah LL, Li Z, Willems van Dijk K, Rensen PCN, Kuivenhoven JA, Groen AK, Hilbers PAJ, van Riel NAW. In vivo and in silico dynamics of the development of Metabolic Syndrome. PLoS Comput Biol. 2018;14(6):e1006145. Epub 20180607. doi: 10.1371/journal.pcbi.1006145. PubMed PMID: 29879115; PMCID: PMC5991635.
- 6. Duarte NC, Becker SA, Jamshidi N, Thiele I, Mo ML, Vo TD, Srivas R, Palsson B. Global reconstruction of the human metabolic network based on genomic and bibliomic data. Proc Natl Acad Sci U S A. 2007;104(6):1777-82. Epub 20070131. doi: 10.1073/pnas.0610772104. PubMed PMID: 17267599; PMCID: PMC1794290.
- 7. Brunk E, Sahoo S, Zielinski DC, Altunkaya A, Dräger A, Mih N, Gatto F, Nilsson A, Preciat Gonzalez GA, Aurich MK, Prlić A, Sastry A, Danielsdottir AD, Heinken A, Noronha A, Rose PW, Burley SK, Fleming RMT, Nielsen J, ..., Palsson BO. Recon3D enables a three-dimensional view of gene variation in human metabolism. Nat Biotechnol. 2018;36(3):272-81. Epub 20180219. doi: 10.1038/nbt.4072. PubMed PMID: 29457794; PMCID: PMC5840010.
- 8. Robinson JL, Kocabaş P, Wang H, Cholley PE, Cook D, Nilsson A, Anton M, Ferreira R, Domenzain I, Billa V, Limeta A, Hedin A, Gustafsson J, Kerkhoven EJ, Svensson LT, Palsson BO, Mardinoglu A, Hansson L, Uhlén M, Nielsen J. An atlas of human metabolism. Sci Signal. 2020;13(624). Epub

- 20200324. doi: 10.1126/scisignal.aaz1482. PubMed PMID: 32209698; PMCID: PMC7331181.
- 9. Fouladiha H, Marashi SA. Biomedical applications of cell- and tissue-specific metabolic network models. J Biomed Inform. 2017;68:35-49. Epub 20170224. doi: 10.1016/j.jbi.2017.02.014. PubMed PMID: 28242343.
- 10. Thiele I, Sahoo S, Heinken A, Hertel J, Heirendt L, Aurich MK, Fleming RM. Personalized whole-body models integrate metabolism, physiology, and the gut microbiome. Mol Syst Biol. 2020;16(5):e8982. doi: 10.15252/msb.20198982. PubMed PMID: 32463598; PMCID: PMC7285886.
- 11. Ben Guebila M, Thiele I. Dynamic flux balance analysis of whole-body metabolism for type 1 diabetes. Nature Computational Science. 2021;1(5):348-61. doi: 10.1038/s43588-021-00074-3.
- 12. Wakil SJ, Abu-Elheiga LA. Fatty acid metabolism: target for metabolic syndrome. J Lipid Res. 2009;50 Suppl(Suppl):S138-43. Epub 20081201. doi: 10.1194/jlr.R800079-JLR200. PubMed PMID: 19047759; PMCID: PMC2674721.
- 13. Fahed G, Aoun L, Bou Zerdan M, Allam S, Bouferraa Y, Assi HI. Metabolic Syndrome: Updates on Pathophysiology and Management in 2021. Int J Mol Sci. 2022;23(2). Epub 20220112. doi: 10.3390/ijms23020786. PubMed PMID: 35054972; PMCID: PMC8775991.
- 14. Qu M, Zhou X, Wang X, Li H. Lipid-induced S-palmitoylation as a Vital Regulator of Cell Signaling and Disease Development. Int J Biol Sci. 2021;17(15):4223-37. Epub 20211011. doi: 10.7150/ijbs.64046. PubMed PMID: 34803494; PMCID: PMC8579454.
- 15. StatPearls. 2022.
- 16. Triplitt CL. Understanding the kidneys' role in blood glucose regulation. Am J Manag Care. 2012;18(1 Suppl):S11-6. PubMed PMID: 22559853.
- 17. Langin D. Adipose tissue lipolysis as a metabolic pathway to define pharmacological strategies against obesity and the metabolic syndrome. Pharmacol Res. 2006;53(6):482-91. Epub 20060327. doi: 10.1016/j.phrs.2006.03.009. PubMed PMID: 16644234.
- 18. Targher G, Bonapace S, Byrne CD. Does high LDL-cholesterol cause cardiovascular disease? Expert Rev Clin Pharmacol. 2019;12(2):91. Epub 20181226. doi: 10.1080/17512433.2019.1561100. PubMed PMID: 30570363.
- 19. Fernandez ML, Jones JJ, Ackerman D, Barona J, Calle M, Comperatore MV, Kim JE, Andersen C, Leite JO, Volek JS, McIntosh M, Kalynych C, Najm W, Lerman RH. Low HDL cholesterol is associated with increased atherogenic lipoproteins and insulin resistance in women classified with metabolic syndrome. Nutr Res Pract. 2010;4(6):492-8. Epub 20101228. doi: 10.4162/nrp.2010.4.6.492. PubMed PMID: 21286407; PMCID: PMC3029790.

- 20. Wang S. Association between serum low-density lipoprotein cholesterol and metabolic syndrome in a working population. Lipids Health Dis. 2021;20(1):73. Epub 20210718. doi: 10.1186/s12944-021-01500-1. PubMed PMID: 34275455; PMCID: PMC8286618.
- 21. van Eunen K, Simons SM, Gerding A, Bleeker A, den Besten G, Touw CM, Houten SM, Groen BK, Krab K, Reijngoud DJ, Bakker BM. Biochemical competition makes fatty-acid β-oxidation vulnerable to substrate overload. PLoS Comput Biol. 2013;9(8):e1003186. Epub 20130815. doi: 10.1371/journal.pcbi.1003186. PubMed PMID: 23966849; PMCID: PMC3744394.
- 22. Panov A, Mayorov VI, Dikalov S. Metabolic Syndrome and β-Oxidation of Long-Chain Fatty Acids in the Brain, Heart, and Kidney Mitochondria. Int J Mol Sci. 2022;23(7). Epub 20220406. doi: 10.3390/ijms23074047. PubMed PMID: 35409406; PMCID: PMC9000033.
- 23. Jensen MD. Fatty acid oxidation in human skeletal muscle. J Clin Invest. 2002;110(11):1607-9. doi: 10.1172/JCI17303. PubMed PMID: 12464664; PMCID: PMC151643.
- 24. Grynberg A, Demaison L. Fatty acid oxidation in the heart. J Cardiovasc Pharmacol. 1996;28 Suppl 1:S11-7. doi: 10.1097/00005344-199600003-00003. PubMed PMID: 8891866.
- 25. Tanaka T, Yamamoto J, Iwasaki S, Asaba H, Hamura H, Ikeda Y, Watanabe M, Magoori K, Ioka RX, Tachibana K, Watanabe Y, Uchiyama Y, Sumi K, Iguchi H, Ito S, Doi T, Hamakubo T, Naito M, Auwerx J, ..., Sakai J. Activation of peroxisome proliferator-activated receptor delta induces fatty acid beta-oxidation in skeletal muscle and attenuates metabolic syndrome. Proc Natl Acad Sci U S A. 2003;100(26):15924-9. Epub 20031215. doi:
- 10.1073/pnas.0306981100. PubMed PMID: 14676330; PMCID: PMC307669.
- 26. Fritzen AM, Lundsgaard AM, Kiens B. Tuning fatty acid oxidation in skeletal muscle with dietary fat and exercise. Nat Rev Endocrinol. 2020;16(12):683-96. Epub 20200922. doi: 10.1038/s41574-020-0405-1. PubMed PMID: 32963340.
- 27. Fisher AB. Intermediary metabolism of the lung. Environ Health Perspect. 1984;55:149-58. doi: 10.1289/ehp.8455149. PubMed PMID: 6376097; PMCID: PMC1568362.
- 28. McKeever TM, Weston PJ, Hubbard R, Fogarty A. Lung function and glucose metabolism: an analysis of data from the Third National Health and Nutrition Examination Survey. Am J Epidemiol. 2005;161(6):546-56. doi: 10.1093/aje/kwi076. PubMed PMID: 15746471.
- 29. Baffi CW, Wood L, Winnica D, Strollo PJ, Gladwin MT, Que LG, Holguin F. Metabolic Syndrome and the Lung. Chest. 2016;149(6):1525-34. Epub

- 20160120. doi: 10.1016/j.chest.2015.12.034. PubMed PMID: 26836925; PMCID: PMC4944780.
- 30. Pillot B, Soty M, Gautier-Stein A, Zitoun C, Mithieux G. Protein feeding promotes redistribution of endogenous glucose production to the kidney and potentiates its suppression by insulin. Endocrinology. 2009;150(2):616-24. Epub 20081009. doi: 10.1210/en.2008-0601. PubMed PMID: 18845639.
- 31. Verma MK, Tripathi M, Singh BK. Dietary Determinants of Metabolic Syndrome: Focus on the Obesity and Metabolic Dysfunction-Associated Steatotic Liver Disease (MASLD). IntechOpen. 2024. Epub 4 Apr. 2024. doi: 10.5772/intechopen.114832.
- 32. Coppack SW, Jensen MD, Miles JM. In vivo regulation of lipolysis in humans. J Lipid Res. 1994;35(2):177-93. PubMed PMID: 8169522.
- 33. Chacko E, Signore C. Five Evidence-Based Lifestyle Habits People With Diabetes Can Use. Clin Diabetes. 2020;38(3):273-84. doi: 10.2337/cd19-0078. PubMed PMID: 32699476; PMCID: PMC7364446.
- 34. Ozen E, Mihaylova R, Weech M, Kinsella S, Lovegrove JA, Jackson KG. Association between dietary saturated fat with cardiovascular disease risk markers and body composition in healthy adults: findings from the cross-sectional BODYCON study. Nutr Metab (Lond). 2022;19(1):15. Epub 20220303. doi: 10.1186/s12986-022-00650-y. PubMed PMID: 35241101; PMCID: PMC8896371.
- 35. Maki KC, Dicklin MR, Kirkpatrick CF. Saturated fats and cardiovascular health: Current evidence and controversies. J Clin Lipidol. 2021;15(6):765-72. Epub 20211001. doi: 10.1016/j.jacl.2021.09.049. PubMed PMID: 34649831.
- 36. Ma Y, Li Y, Chiriboga DE, Olendzki BC, Hebert JR, Li W, Leung K, Hafner AR, Ockene IS. Association between carbohydrate intake and serum lipids. J Am Coll Nutr. 2006;25(2):155-63. doi: 10.1080/07315724.2006.10719527. PubMed PMID: 16582033; PMCID: PMC1479303.
- 37. Jian ZH, Chiang YC, Lung CC, Ho CC, Ko PC, Ndi Nfor O, Chang HC, Liaw YC, Liang YC, Liaw YP. Vegetarian diet and cholesterol and TAG levels by gender. Public Health Nutr. 2015;18(4):721-6. Epub 20140625. doi: 10.1017/S1368980014000883. PubMed PMID: 24963684; PMCID: PMC10271072.
- 38. Leonhardt M, Langhans W. Fatty acid oxidation and control of food intake. Physiol Behav. 2004;83(4):645-51. doi: 10.1016/j.physbeh.2004.07.033. PubMed PMID: 15621070.
- 39. Paoli A, Bosco G, Camporesi EM, Mangar D. Ketosis, ketogenic diet and food intake control: a complex relationship. Front Psychol. 2015;6:27. Epub 20150202. doi: 10.3389/fpsyg.2015.00027. PubMed PMID: 25698989; PMCID: PMC4313585.

- 40. Abbasian M, Delvarianzadeh M, Ebrahimi H, Khosravi F. Lipid ratio as a suitable tool to identify individuals with MetS risk: A case- control study. Diabetes Metab Syndr. 2017;11 Suppl 1:S15-S9. Epub 20160822. doi: 10.1016/j.dsx.2016.08.011. PubMed PMID: 27575046.
- 41. Yasin A, Nguyen M, Sidhu A, Majety P, Spitz J, Asgharpour A, Siddiqui MS, Sperling LS, Quyyumi AA, Mehta A. Liver and cardiovascular disease outcomes in metabolic syndrome and diabetic populations: Bi-directional opportunities to multiply preventive strategies. Diabetes Res Clin Pract. 2024;211:111650. Epub 20240409. doi: 10.1016/j.diabres.2024.111650. PubMed PMID: 38604447.
- 42. Cantley J. The control of insulin secretion by adipokines: current evidence for adipocyte-beta cell endocrine signalling in metabolic homeostasis. Mamm Genome. 2014;25(9-10):442-54. Epub 20140822. doi: 10.1007/s00335-014-9538-7. PubMed PMID: 25146550.
- 43. Santoro A, Kahn BB. Adipocyte Regulation of Insulin Sensitivity and the Risk of Type 2 Diabetes. N Engl J Med. 2023;388(22):2071-85. doi: 10.1056/NEJMra2216691. PubMed PMID: 37256977.
- 44. Koff SA. Guidelines to determine the size and shape of intestinal segments used for reconstruction. J Urol. 1988;140(5 Pt 2):1150-1. doi: 10.1016/s0022-5347(17)41985-9. PubMed PMID: 3184288.
- 45. Ordovas JM, Ferguson LR, Tai ES, Mathers JC. Personalised nutrition and health. BMJ. 2018;361:bmj.k2173. Epub 20180613. doi: 10.1136/bmj.k2173. PubMed PMID: 29898881; PMCID: PMC6081996.
- 46. Noronha A, Modamio J, Jarosz Y, Guerard E, Sompairac N, Preciat G, Daníelsdóttir AD, Krecke M, Merten D, Haraldsdóttir HS, Heinken A, Heirendt L, Magnúsdóttir S, Ravcheev DA, Sahoo S, Gawron P, Friscioni L, Garcia B, Prendergast M, ..., Thiele I. The Virtual Metabolic Human database: integrating human and gut microbiome metabolism with nutrition and disease. Nucleic Acids Res. 2019;47(D1):D614-D24. doi: 10.1093/nar/gky992. PubMed PMID: 30371894; PMCID: PMC6323901.
- 47. McCreery CV, Alessi D, Mollo K, Fasano A, Zomorrodi AR. Investigating intestinal epithelium metabolic dysfunction in Celiac Disease using personalized genome-scale models. bioRxiv. 2024:2024.04.29.591234. doi: 10.1101/2024.04.29.591234.
- 48. Sahoo S, Thiele I. Predicting the impact of diet and enzymopathies on human small intestinal epithelial cells. Hum Mol Genet. 2013;22(13):2705-22. Epub 2013/03/13. doi: 10.1093/hmg/ddt119. PubMed PMID: 23492669; PMCID: PMC3674809.
- 49. Heirendt L, Arreckx S, Pfau T, Mendoza SN, Richelle A, Heinken A, Haraldsdóttir HS, Wachowiak J, Keating SM, Vlasov V, Magnusdóttir S, Ng CY,

Preciat G, Žagare A, Chan SHJ, Aurich MK, Clancy CM, Modamio J, Sauls JT, ..., Fleming RMT. Creation and analysis of biochemical constraint-based models using the COBRA Toolbox v.3.0.

Supplementary files

Supplementary Table 1. List of reactions related to the MetS biomarkers and fatty acid beta-oxidation pathway within the male and female WBMs, along with their predicted flux values under the examined diets.