Impossible to go Beyond Beef? A Nutriomics Comparison

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1 Abstract

2 Concerns regarding the effects of red meat on human and environmental health are prompting 3 consumer interest in plant-based diets. As global food systems strive to meet the dietary needs of 4 an estimated mid-century population of 10 billion, a new generation of plant-based meat 5 alternatives—formulated to mimic the taste and nutritional composition of red meat—have 6 attracted considerable consumer interest, research attention, and media coverage. We used 7 untargeted metabolomics to provide an in-depth comparison of the nutrient profiles of grass-fed 8 ground beef and a market-leading plant-based meat alternative. Metabolomics revealed a 90% 9 difference in nutritional profiles beef and a popular plant-based meat, many of which can have 10 important consumer health implications. This information could not be determined from their 11 Nutrition Facts, which suggests nutritional similarity. Our findings indicate that beef and a



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- 12 popular plant-based meat should not be viewed as nutritionally interchangeable, but as
- 13 complementary in terms of provided nutritional entities. As society aims to increase food
- 14 production with ~ 60% by 2050, the meat and the plant-based meat industries will likely coexist
- 15 and have to complement each other in order this reach this goal.

16 **Main**

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value of \$1142.9 billion by 2023^5 .

17 By 2050, global food systems will need to meet the dietary demands of almost 10 billion 18 people. To meet these demands in a healthy and sustainable manner, it is suggested that diets 19 would benefit from a shift towards consumption of more plant-based foods and less meat, particularly in Western countries¹. This has raised questions of whether novel plant-based meat 20 21 alternatives represent healthy and sustainable alternatives to meat^{2,3}. The new generation of plant-based meats such as the ImpossibleTM Burger and Beyond 22 Burger[®] are becoming increasingly popular with consumers. Their success has led other 23 24 international food companies—including traditional meat companies—such as Purdue Farms 25 (US), Cargill (US), Lightlife (US), Gardein Protein International (Canada), Maple Leafs

(Canada), Quorn (UK), Tyson Foods (US), and Unilever (UK/The Netherlands) to invest in their
own versions of these products⁴. The global plant-based meat sector is currently experiencing
rapid growth and is projected to increase from \$11.6 billion in 2019 to \$30.9 billion by 2026⁵
with a compound annual growth rate (CAGR) of 15% (Fig. 1). In contrast, the animal meat
sector is "only" expecting a CAGR of 3.9% during that time (Fig. 1) and will reach a market

The production of plant-based meats as a replacement for animal-sourced meat is nothing new. One of the earliest engineered meat alternatives was ProtoseTM, a plant-based meat made from wheat gluten, peanuts and, soybean oil, which was designed by John Kellogg in the late nineteenth century. In 1899, Kellogg wrote the following in his patent application for ProtoseTM: *"The objective of my invention is to furnish a vegetable substitute for meat which shall possess equal or greater nutritive value in equal or more favorable form for digestion and assimilation and which shall contain the essential nutritive elements in approximately the same* 39 proportion as beef and mutton and which substitute has a similar flavor and is as easily
40 digestible as the most tender meat" (U.S. Patent No 670283A).

41

41 Unlike previous products, contemporary plant-based meat alternatives have accomplished 42 to create a taste and sensory experience that more closely resembles red meat. For example, soy 43 leghemoglobin imitates the "bloody" appearance and taste of heme proteins in meat, while 44 extracts from red beets, red berries, carrots, and/or other similarly colored vegetables are often 45 embedded in plant-based products to give them a reddish 'meat-like' appearance⁴. Methyl 46 cellulose is often used to give plant-based meat alternative a 'meat-like' texture. Modern meat 47 alternatives also match the protein content of meat by using isolated plant proteins (e.g., soy, pea, 48 potato, mung bean, rice, mycoprotein, and/or wheat) and they are often fortified with vitamins 49 and minerals naturally found in red meat (e.g., vitamins B_{12} , zinc, and iron) to provide an even 50 more direct nutritional replacement⁶. Indeed, a popular novel soy-based alternative closely 51 matches the Nutrition Facts panel of beef (Fig. 2), and to consumers reading nutritional labels they appear nutritionally interchangeable⁷. Nonetheless, food sources in their natural state have 52 53 considerable complexity and contain a wide variety of nutrients (e.g., phenols, anti-oxidants, 54 peptides, amino acids, fatty acids, carboxylic acid etc.), the majority of which do not appear on nutrition labels⁸, but have important health implications. Important nutritional differences are 55 likely to exist between beef and the new generation of plant-based meat replacements; however, 56 57 this has not been thoroughly assessed.

Given the scientific and commercial interest in plant-based meat alternatives, the goal of our study was to use untargeted metabolomics to provide an in-depth comparison of the nutrients in grass-fed ground beef and a popular next-gen soy-based meat alternative, both of which may be considered healthier and more environmentally friendly sources of "beef"^{4,9}. Metabolomics is an analytical profiling technique that allows researchers to measure and compare large numbers
of nutrients and metabolites that are present in biological samples. Metabolomics analysis
enabled a look "behind the curtain" to evaluate how beef and a popular soy-based alternative
differ nutritionally—beyond what their labels reveal (Fig. 2).

66

67 Untargeted Metabolomics of Plant-Based Meat and Beef

68 A schematic representation of the study flow is provided in Fig. 2. We purchased 69 eighteen packages of a popular next-gen soy-based meat alternative from a local grocery store. 70 Ground beef from eighteen grass-fed cattle was purchased from Alderspring Ranch (May, ID) 71 and matched for fat (14 grams) and serving size (113 grams) to the soy-based alternative. To 72 identify potential nutritional differences between beef and the soy-based meat alternative, we 73 analyzed the relative abundance of metabolites in individually cooked samples (n=18 beef 74 samples and n=18 sov-based meat alternative samples, respectively) using gas 75 chromatography/electron-ionization mass spectrometry (GC/ei-MS)-based untargeted metabolomics¹⁰. We profiled 190 unique metabolites in the beef and soy-based meat samples, 76 77 which were tested for differences between products using the Wilcoxon rank sum test with 78 Benjamini-Hochberg adjusted *P*-values at 5% (False Discovery Rate; FDR < 0.05). 79 We found that a total of 171 out of 190 profiled metabolites (90%) were different (FDR <80 0.05) between beef and the soy-based alternative (Table S1). To visualize differences and 81 identify the top metabolites that contributed to the nutritional disparity between beef and plant-82 based meat, we created a ranked heatmap of the top fifty metabolites based on the Pearson 83 distance measure and the Ward clustering algorithm, and performed unsupervised principal 84 component analysis using software procedures from MetaboAnalyst 4.0

85	(http://www.metaboanalyst.ca). Both the heatmap (Fig. 3A) and unsupervised principal
86	component analysis (Fig. 3B) revealed a distinct separation in nutritional components between
87	the grass-fed ground beef and the soy-based meat alternative. To identify the main nutrient
88	classes that differed between beef and the soy-based alternative, we then clustered individual
89	metabolites into nutrient classes according to their structural similarity using Chemical Similarity
90	Enrichment Analysis (ChemRICH) software procedures (<u>http://chemrich.fiehnlab.ucdavis.edu/</u>).
91	We identified 24 nutrient classes with \geq 3 structurally similar metabolites regardless of
92	whether these metabolites were found in beef or the plant-based meat (Table 1). We found that
93	23 of the nutrient classes differed significantly (FDR < 0.05) between beef and the soy-based
94	meat alternative (Table 1). Several nutrients were found either exclusively (22 metabolites total)
95	or in greater quantities in beef (52 metabolites total) compared with the soy-based meat
96	alternative (Table S1). Similarly, several other nutrients were found exclusively (31 metabolites
97	total) or in greater quantities (67 metabolites total) in the soy-based meat alternative when
98	compared to beef.
99	Creatinine (product of creatine), hydroxyproline (a non-proteinogenic amino acid),
100	anserine (a carnosine metabolite), glucosamine (a saccharide), and cysteamine (an aminothiol)
101	are examples of nutrients only found in beef and appeared as discriminating metabolites within
102	their respective nutrient class (Table 1). These nutrients have important physiological, anti-
103	inflammatory, and/or immunomodulatory roles ^{11,12} and low intakes are associated with
104	cardiovascular, neurocognitive, retinal, hepatic, skeletal muscle, and connective tissue
105	dysfunction ^{11,12} . For example, creatine and anserine provide neurocognitive protection in older
106	adults ^{13,14} . Cysteamine, a potent antioxidant, also has neuroprotective effects and is a precursor
107	of glutathione—one of the most potent intracellular antioxidants ¹⁵ . Squalene has strong anti-

oxidant, anti-bacterial, and anti-tumor activity¹⁶, while dietary hydroxyproline and glucosamine
 stimulate collagen biosynthesis and are important for maintaining the structure and strength of
 connective tissue and blood vessels^{11,17}.

111 On the other hand, metabolites in nutrient classes such as phenols, tocopherols, and 112 phytosterols (Table 1) were found exclusively or in much greater abundance in the plant-based 113 meat when compared to beef. For instance, the plant-based meat alternative contained more 114 tocopherols (α , γ , and δ)—a class of nutrients with vitamin E activity best known for their antioxidant effects¹⁸. We also found several phytosterols such as *beta*-sitosterol, campesterol, 115 116 and stigmasterol in the plant-based meat, which collectively possess antioxidant, antiinflammatory, and cancer-protective properties¹⁹. We also found a wider variety and greater 117 118 abundance of phenolic compounds in the soy-based alternative when compared to beef (Table 1). 119 Identified compounds include sulfurol, syringic acid, vanillic acid, and methylated/hydroxylated 120 forms of valeric acid, which can benefit human health by dampening oxidative stress and inflammation²⁰. 121 122 Within the nutrient class of polyunsaturated fatty acids (PUFAs); arachidonic acid (ARA, 123 C20:4, ω -6) and docosahexaenoic acid (DHA, C22:6, ω -3) were found exclusively (DHA) or in 124 much greater quantities (ARA) in the grass-fed beef samples (Table 1). These essential fatty 125 acids are major constituents of the brain phospholipid membrane and have important roles in

cognition, immunomodulation, platelet function, and cell signaling^{12,21}. Their deficiencies are
associated with cognitive decline and increased risk of cardiovascular disease^{12,21}.

Important differences were also observed in saturated fatty acid and glyceride classes
(Table 1). The main saturated fatty acids and glycerides (Table 1) in the plant-based meat were
coconut oil-derived lauric acid, monolaurin, dilaurin, and trilaurin, which possess anti-microbial

and/or anti-inflammatory properties²². On the other hand, we found higher levels of the dietary
odd-chain saturated fatty acids (OCFAs) pentadecanoic acid (C15:0) and heptadecanoic acid
(C17:0) in beef than in the soy-based alternative. These compounds are believed to exert their
beneficial effects by attenuating inflammation, dyslipidemia, and cell fibrosis²³, and increased
dietary intake is associated with a lower risk of metabolic disease^{24,25}.

For an exhaustive list of the different metabolites found in beef and the plant-based meat and their potential roles in human health, the readers are referred to Table S1. While several of these nutrients are considered non-essential or conditionally-essential based on life-stages (*e.g.*, infancy, pregnancy, or advanced age) and are often less appreciated in discussions of human nutritional requirements⁸, their importance should not be ignored as low intakes can have profound impacts on human health.

142

143 Can Plant-Based Meat Alternatives Meet Human Nutritional

144 **Requirements?**

145 A key question in the broader discussion of replacing of animal foods with plant-based 146 substitutes is whether plant-based substitutes can adequately satisfy human nutrition requirements. The underlying dietary strategy for most of mankind now²⁶, and certainly throughout our 147 evolutionary history, has been omnivory^{27,28}. While overlap exists between nutritional profiles of 148 149 animal and plant foods, needs for certain nutrients—including vitamins C and E (tocopherols), 150 folate, manganese, thiamin (B₁), potassium, phenols, and other phytochemicals—are more 151 readily met by consuming plant foods. However, needs for other nutrients—including heme-iron, 152 retinol (vitamin A), vitamin B₁₂, and long-chain PUFAs, and secondary nutrients such as 153 creatine, anserine, taurine, and cysteamine—are met more readily or exclusively from animal

foods. Animal foods also facilitate uptake of several plant nutrients (*e.g.*, non-heme iron and
zinc)^{29,30}, while plant nutrients (*e.g.*, phytochemicals and fiber) provide protective effects against
potentially harmful compounds (*e.g.*, heterocyclic amines, advanced glycation end products etc.)
in cooked and cured animal foods³¹. The secondary compounds in plant foods (*i.e.*,
phytochemicals) also exert key antioxidant, anti-inflammatory, anticancer, and
immunomodulatory roles³². Arguably, plant and animal foods in the human diet interact
symbiotically to improve human health.

161 Nonetheless, those following vegan and vegetarian diets often have improved metabolic 162 health when compared to omnivores, though differences may disappear when extensively adjusting for lifestyle and dietary factors^{33,34}. For example, large-scale population based studies 163 performed in individuals with 'healthy lifestyles' such the Oxford-EPIC Study³⁵ (n~64,000) and 164 the 45-and-Up Study $(n\sim 267,000)^{36}$ report no difference in mortality rates between omnivores 165 166 and vegetarians, when omnivores also consume high amounts of fruits, vegetables, nuts, and seeds. Additionally, intra-individual differences in nutrient metabolism³⁷⁻⁴⁰ may explain why 167 168 some individuals can thrive on plant-based diets, while others experience health problems associated with nutrient deficiencies⁴¹. While discussions regarding red meat, plant-based diets, 169 and human health have become increasingly vigorous in recent times^{42,43}, academics^{44,45} and 170 171 governing bodies⁴⁶ generally agree that population health, particularly in Western countries, 172 would benefit from a shift towards increasing the amount of whole food plant-sources as opposed to consuming a Standard American/Western diet—rich in ultra-processed foods^{47,48}. 173 174 While plant-based foods are often considered to be healthy foods to consume, Hu and colleagues² have expressed concern in extending these notions to plant-based meat alternatives 175 176 given their ultra-processed nature. Of note is a recent 8-week randomized controlled trial (RCT)

177 that compared biomarkers of metabolic health in response to consumption of ~ 2.5 servings/day of a market leading plant-based alternative (Beyond MeatTM) versus organic animal meats (grass-178 fed beef, organic chicken, and pork), both consumed as part of an omnivorous diet⁴⁹. The authors 179 180 found that serum trimethylamine-N-oxide (TMAO) concentrations were lower following 8 181 weeks of plant-based meat consumption when compared to animal meats, but only if the 182 participants received the plant-based meat intervention first. Participants in the plant-based meat 183 arm also lost weight when compared to the animal-based group, but again only if the plant-based 184 meats were consumed first, not second. No order effect was observed for low density 185 lipoprotein-cholesterol (LDL-C), which was lower after plant-based meat ingestion regardless of 186 the order of intervention. No group differences were observed in other health biomarkers (high 187 density lipoprotein-cholesterol, triglycerides, insulin, glucose and blood pressure). 188 TMAO is a gut microbiota-dependent metabolite produced from quaternary ammonium 189 compounds such as phosphatidylcholine, choline, betaine, and L-carnitine, which are 190 predominantly found in animal meats, but TMAO can also be directly obtained from seafood⁵⁰. 191 Whether TMAO is truly an effector of metabolic disease in otherwise healthy individuals and 192 whether increased TMAO levels in cardiovascular disease and type 2 diabetes is the result (rather than the cause) of disease-related dysbiosis is currently a focal point of discussion^{50,51}, and likely 193 194 depends on the context in which elevated TMAO levels are observed (pathophysiological states versus dietary intakes of fish and red meat as part of an otherwise "healthy diet")⁵⁰. Nonetheless, 195 196 this work provides preliminary evidence that a "flexitarian approach" (replacing some meat with 197 plant-based alternatives as part of an omnivorous diet) has no negative health effects and may have slight positive benefits in terms of weight control and cardiometabolic risk profiles⁴⁹. 198 199 Future work that assesses additional health biomarkers (e.g., disease-associated inflammation

and oxidative stress) and is aimed at elucidating mechanistic pathways by which plant-based
 meat alternatives impact metabolic health are needed to confirm potential health effects of plant based meat alternatives.

203 Similarities between beef and the soy-based alternative in terms of total protein content 204 and several vitamins and minerals (Fig. 2.) suggests that a "flexitarian approach" (replacing 205 some meat with plant-based alternatives as part of an omnivorous diet) is unlikely to negatively 206 impact nutritional status of consumers in the long-run, but this also depends on what other foods 207 are part of the diet and the degree to which plant-based substitutes replace animal foods (e.g., the 208 occasional replacement or full replacement of all animal foods). If a particular nutrient is 209 obtained in sufficient quantities from other commonly consumed foods then its lack in a plantbased meat is likely of no consequence⁴⁹. However, caution is warranted for vulnerable 210 211 populations such as children, women of childbearing age, and older individuals who may be at increased risk for nutritional deficiencies with low intakes of animal foods^{52,53}. Moreover, in 212 213 discussions about replacing meat with plant-based substitutes on a global level, it is important 214 that food policies do not adversely impact the estimated 2 billion people in developing countries whose basic nutritional needs and livelihoods depend on meat and livestock products^{3,52}. 215 216 Our work has several limitations. While the soy-based meat alternative we studied is one 217 of the most popular products currently on the market, product formulations of next-gen plant-

based meats differ slightly in terms of the type of isolated plant proteins (*e.g.*, soy, pea, potato,

219 mung bean, rice, mycoprotein and/or wheat), fats (*e.g.*, canola, soy coconut, and/or sunflower

220 oil), and/or other ingredients (*e.g.*, soy leghemoglobin, different vegetable extracts, and/or

221 different flavoring agents)⁶. Nonetheless, we reasonably expect that plant-based meat alternatives

are far more similar to each other than they are to red meat.

The nutritional components highlighted in our work represent only a small fraction of the currently estimated >4,000 distinct metabolites present in foods such as beef and soy (the main constituent of the studied plant-based meat alternative)⁵⁴—many of which have known health effects, but would require extensive targeted metabolomics approaches for their systematic identification.

As the field of nutriomics (the application of metabolomics in nutrition domains) progresses, we will undoubtedly gain greater appreciation of the complexity of natural food matrices and the ability of manifold nutritional constituents to synergistically modulate human health⁸. The complexity of the natural food matrix highlights that attempting to mimic natural food sources using single constituents such as isolated proteins, vitamins, and minerals is challenging and underestimates the true nutritional complexity of food sources in their natural state.

235

236 **Conclusions**

237 Untargeted metabolomics revealed a 90% difference in nutritional profiles between beef 238 and a market-leading soy-based meat alternative. This information could not be determined from 239 their Nutrition Facts panels (Fig. 2.), which suggests that similar nutrients can be obtained from 240 both products. While beef and the soy-based alternative both contain a wide range of potentially 241 beneficial nutrients (e.g., phenols, tocopherols, fatty acids, antioxidants, amino acids, and 242 dipeptides) as well as some potentially deleterious compounds (e.g., maillard reaction end-243 products) (Table 1 and Table S1), large differences in individual nutrients indicate that these 244 products should not be viewed as nutritionally interchangeable (Fig. 3 and Table S1). This information does not appear to be known with consumers⁷. Thus, the new information we 245

provide is important for making informed decisions by consumer decisions and to inform food
policies and dietary advice. It cannot be determined from our data if either source is healthier to
consume.

249 As society strives to meet dietary needs of an estimated 10 billion people by 2050, the 250 challenge is to create global food systems that are locally adapted to meet dietary needs in a 251 sustainable, healthy, and inclusive manner³. Animal and plant foods—and the nutrients they 252 provide—should arguably be viewed as complementary rather than competitive in this scenario. 253 The observed nutritional differences between beef and a popular plant-based meat alternative 254 further highlights this notion. As global food systems work to increase production with $\sim 60\%$ by 255 2050, both the meat and plant-based alternative industries will likely coexist and have to 256 complement each other in order to meet this lofty goal³.

257

258 Methods

259 **Product sourcing**

260 Eighteen different packages (340 grams or 12 oz each) of a market-leading plant-based 261 meat alternative was bought from a local grocery store in Raleigh, NC, USA. Ground beef from 262 eighteen grass-fed, black angus cattle (454 grams or 16 oz each) was purchased from 263 Alderspring Ranch (May, ID) and matched for total fat content (14 grams) to the soy-based 264 alternative, which was confirmed using proximate analysis (method AOAC 960.39; Microbac 265 Laboratories, Warrendale, PA). Individual patties (112 grams or 4 oz each) were formed from 266 each individual package of plant-based meat and beef, respectively. Individual patties were 267 cooked on a non-stick skillet until the internal temperature of each patty read 71 °C as 268 determined by a meat thermometer. One-gram microcore samples were obtained from the middle of each patty (n=18 for ground beef; n=18 for soy-based meat replacement) using a bioptome
device, immediately frozen in liquid nitrogen, and stored at -80 degrees °C until metabolomics
analysis.

272

273 Sample preparation

274 Microcore samples the plant-based meat replacement and bovine skeletal muscle (*i.e.*, 275 beef) were powdered under liquid N₂ and homogenized in 50% aqueous acetonitrile containing 276 0.3% formic acid (50 mg wet weight sample per ml homogenate) using a Qiagen Retsch Tissue 277 Lyser II set to a frequency of 30 oscillations/sec for a total of 2 min with one 5 mm glass ball 278 (GlenMills, Inc, #7200-005000TM) per tube. 100 µl of each sample homogenate was then transferred into a fresh, 1.5-ml, Reduced Surface Activity (RSATM) glass autosampler vial 279 280 (catalog number 9512C-1MP-RS, MicroSolv Technology Corporation, Leland, NC). Proteins in 281 sample homogenates were subsequently "crash" precipitated with 750 µl dry methanol spiked 282 with C14:0-D₂₇ (perdeuterated myristic acid, Sigma 366889, 6.25 mg/liter, CN167: 141; CN188: 283 115) and centrifuged at 13.500 x g rcf for 5 minutes (Vial CentrifugeTM, MicroSolv, catalog 284 C2417). The crash solvent is spiked with with C14:0- D_{27} Myristic Acid as an internal standard 285 for retention-time locking (described below). 700 µl of the supernatant of each sample 286 homogenate were subsequently transfered to fresh RSATM glass vials (catalog number 9512C-287 1MP-RS, MicroSolv Technology Corporation, Leland, NC). Methanolic extracts were then dried 288 in a Savant SPD111V SpeedVac Concentrator (Thermo Scientific, Asheville, NC), with the help 289 of a final pulse of toluene (Fisher Scientific, catalog number T324-50) as an azeotropic drying 290 agent. 25 µl methoxyamine hydrochloride (18 mg/ml in dry pyridine: Fisher Scientific, catalog 291 number T324-50) was then added to each sample and incubated at 50 °C for 30 minutes for

292 methoximation of certain reactive carbonyl groups. Finally, metabolites were rendered volatile

by replacement of easily exchangeable protons with trimethylsilyl (TMS) groups using N-

294 methyl-*N*-(trimethylsilyl) trifluoroacetamide (MSTFA; 75 µl per sample Cerilliant M-132,

- 295 Sigma, St. Louis, MO) at 50 °C for 30 minutes.
- 296

297 (GC/ei-MS) analysis

298 Samples were run on a 7890B GC / 5977B single-quadrupole, Inert MS (Agilent 299 Technologies, Santa Clara, CA). This system is equipped with a MultiMode Inlet, which, in 300 combination with a mid-column, purged ultimate union (PUU), enables hot back-flushing of the 301 upstream half of the column at the end of each run to reduce fouling of both GC and MS with 302 heavy contaminants ("high boilers") and carryover between injections. Briefly, the two wall-303 coated, open-tubular (WCOT) GC columns connected in series are both from J&W/Agilent (part 304 122-5512 UI), DB5-MS UI, 15 meters in length, 0.25 mm in diameter, with a 0.25-µm luminal 305 film. This film is a nonpolar, thermally stable, phenyl-arylene polymer, similar in performance 306 to traditional 5%-phenyl-methylpolysiloxane films. Prior to each daily run, the starting inlet 307 pressure is empirically adjusted such that the retention time of the TMS-D27-C14:0 standard is 308 set at ~16.727 minutes. After a quick, initial distillation within the MMI, the GC oven ramps 309 from 60-325 °C at a speed of 10 °C/minute. Under these conditions, derivatized metabolites 310 elute from the column and reach the MS detector at known times (e.g., bis-TMS-lactic acid at 311 ~6.85 minutes, and TMS-cholesterol at ~27.38 minutes). A mid-column pneumatic device (PUU) 312 provides a means for hot back-flushing of the upstream GC column at the end of each run while 313 the oven is held at 325 °C for a terminal "bake-out" as an antifouling and anti-carryover measure 314 (analogous to that devised by Chen et al. 2009). During this terminal "bake-out," the inlet is also

315 held at 325 °C while it is purged via its split-flow, waste vent with a large flow of the carrier gas, 316 helium. Radical cations generated with conventional electron ionization via a tungsten-rhenium 317 filament set to an energy of 70 eV are scanned broadly from 600 to 50 m/z in the detector 318 throughout the run. Cycle time is approximately 38 minutes. We typically derivatize and run 319 daily batches of ~28 unknowns and a processed blank ("ghost" sample). Our GC/MS methods are based on validated methods and generally follow those of Roessner et al. (2000)⁵⁵, Fiehn et 320 al. (2008)⁵⁶, Kind et al. (2009)⁵⁷, McNulty et al. (2011)⁵⁸, Banerjee et al. (2015)⁵⁹, and Clinton et 321 al. $(2020)^{60}$. 322

323

Data reduction

325 Raw data from Agilent's MassHunter software environment were imported into the 326 freeware, Automatic Mass Spectral Deconvolution and Identification Software or AMDIS 327 (version 2.73), developed by Drs. Steve Stein, W. Gary Mallard, and their coworkers at National Institute of Standards and Technology or NIST (Mallard and Reed 1997⁶¹, Halket et al. 1999⁶², 328 Stein 199963; courtesy of NIST at http://chemdata.nist.gov/mass-spc/amdis/). Deconvoluted 329 330 spectra were annotated as metabolites, to the extent possible, using an orthogonal approach that 331 incorporates both retention time (RT) from GC and the fragmentation pattern observed in EI-MS, 332 both of which can be remarkably reproducible with contemporary instrumentation. Peak 333 annotation was based primarily on our own RT-locked spectral library of metabolites (2059 334 spectra from 1174 unique compounds, and growing). Our library is built upon the Fiehn GC/MS 335 Metabolomics RTL Library (a gift from Agilent, their part number G1676-90000; Kind et al. 2009⁵⁷. Additional spectra have been gleaned from running pure reagent standards in our lab, 336 337 from the Golm Metabolome Library (courtesy of Dr. Joachim Kopka and coworkers at the Max

338 Planck Institute of Molecular Plant Physiology, Golm, Germany; Kopka *et al.* 2005⁶⁴;

339 <u>http://csbdb.mpimp-golm.mpg.de/csbdb/gmd/gmd.html</u>), and from the Wiley 10th-NIST 2014

340 commercial library (Agilent G1730-64000). Peak alignment and chemometrics of log-base-two-

341 transformed areas of deconvoluted peaks were performed with our own custom macros, written

in our lab in Visual Basic (version 6.0) for use in the Excel (Microsoft Office Professional Plus

343 2019) software environment (both from Microsoft, Redmond, WA). The full list of annotated

344 metabolites and their retention times presented in Table S2.

345

346 Data processing

347 Three investigators (SVV, JRB, and MJM) subsequently performed line-by-line manual 348 curation to fix miscalls and highlighted ambiguities inherent in certain isomeric or otherwise 349 similar metabolites. Metabolites were retained for further analysis if detected in $\geq 80\%$ of 350 samples of either the plant-based meat replacement or ground beef (*i.e.*, 14 out of 18 samples per 351 group). If Th. As can be observed from Table S1, this was the case for 53 metabolites, which 352 were related detected in one source (e.g., beef or plant-based alternative) but not the other. A 353 total of 31 metabolites were detected only on the plant-based meat samples but remained absent 354 in all beef samples; while 22 metabolites were found in beef samples but remained absent in the 355 plant-based meat. In the case of remaining missing values in other metabolites-for which a 356 signal was detected in \geq 14 out of 18 samples in one group (beef or plant) and \geq 1 sample of the other group—k-nearest neighbor imputation was performed^{65,66}. 357

This decision was made after careful deliberation with colleagues at the Biostatistics and the Metabolomics Core at Duke University, and was based on the expectation that in such cases the metabolite feature was truly nonexistent (or at least below the Level of Detection) for a given group (beef or plant meat) and was not due to chromatographic non-detection. In other words, 362 had the metabolite been present in the food source at meaningful levels, it would have registered 363 as we detected this metabolite in \geq 80% of samples in the other group (*i.e.*, 14 out of 18 samples). 364 To illustrate this with an example; anserine (β -alanyl-1-methyl-l-histidine; a methylated 365 product of carnosine) is metabolite that is well-known to occur in beef and other animal meats, but known to be absent in plant samples¹¹. Similarly, soy isoflavones such as β -sitosterol and 366 367 campasterol would normally not be found in grass-fed beef, but were readily detected in all 368 plant-based meat samples (Fig. S2.). If we used KNN imputation (or other commonly used 369 imputation methods such as PLS, SVD, BPCA etc.) without accounting for true absence of 370 metabolites in a given group, our data set would falsely imply that some metabolites are in the 371 plant or beef source of which we know with certainty that they cannot be there, which we argue 372 would be incorrect to report.

373

374 Data analysis

375 After data processing, individual metabolites were tested for normality using 376 Kolmogorov-Smirnov tests (p < 0.05) using SAS 9.4 (Cary, North Carolina, USA). Several 377 metabolites did not show a normal distribution after log transformation, which may be expected 378 based on the large differences between beef and the plant-based meat alternative—53 379 metabolites were detected exclusively in only either the plant-based meat or beef and had log-380 transformed values close to 0. To test differences in individual metabolites between groups, we 381 subsequently used the non-parametric Wilcoxon with Benjamini-Hochberg adjusted p-values at 5% to account for false discovery (FDR < 0.05). 382 383 Bioactivities and potential health effects of annotated metabolites were explored by

384 entering Chemical Abstracts Service (CAS) # of individual metabolites in FooDB

385 (https://foodb.ca/) and/or PubChem (https://pubchem.ncbi.nlm.nih.gov/) databases, while 386 metabolic pathway identification of individual metabolites was performed using the Kyoto 387 Encyclopedia of Genes and Genomes (KEGG) (https://www.genome.jp/). To inform the 388 discussion of metabolomics findings, we clustered metabolites by chemical class using freely-389 available ChemRICH software procedures (http://chemrich.fiehnlab.ucdavis.edu/; courtesy of 390 Dr. Oliver Fiehn and coworkers at the University of California, Davis, USA⁶⁷ (Fig. S2.). To 391 enable cluster analysis via structural similarity and ontology mapping, InChiKeys, PubChemID 392 and SMILES canonicals for each metabolite was retrieved by entering its respective Chemical 393 Abstracts Service (CAS) # in the PubChem (https://pubchem.ncbi.nlm.nih.gov/). After 394 ChemRICH analysis, investigators performed line-by-line manual curation to fix any apparent 395 miscalls or apparent misclassification of individual metabolites and to perform manual 396 adjustment of metabolite classification when appropriate (e.g., ChemRICH classified pyrodixine 397 as a separate "Vitamin B6" category in which case the metabolite was lumped into a larger class 398 simply named "Vitamins"), after which analysis was re-ran. Finally, to visualize differences in 399 individual metabolites between groups and identify the top metabolites that contributed to the 400 nutritional differences between beef and the plant-based meat replacement, we created a ranked 401 heatmap of the top fifty metabolites based on the Pearson distance measure and the Ward 402 clustering algorithm and performed unsupervised principal component analysis using software 403 procedures from MetaboAnalyst 4.0 (https://www.metaboanalyst.ca) (Fig. 3).

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604

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612	consumes a vegetarian diet.
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627 **Contributions**

628 S.V.V., F.D.P., and S.L.K contributed to the conception and design of the study. S.V.V., J.R.B.,

- and M.J.M. were responsible for the metabolomics analysis of the study. S.V.V., C.F.P., and
- 630 K.M.H. performed the statistics. S.V.V and F.D.P. drafted the manuscript and all authors
- 631 contributed to critical revisions of the manuscript for important intellectual content. S.V.V. had
- full access to the data and takes responsibility for the integrity of the data and the accuracy of the
- data analysis; S.V.V. affirms that the manuscript is an honest, accurate, and transparent account
- of the study being reported; that no important aspects of the study have been omitted; and that
- 635 any discrepancies in the analysis have been explained.
- 636

637 Competing Interests

- 638 The authors declare no competing interests.
- 639

640 **Data and materials availability**.

- 641 All data that support the findings of this study are available in the main text, tables/figures,
- and/or the supplementary materials. The full metabolomics data set is available at Dryad:
- 643 <u>https://doi.org/10.5061/dryad.3ffbg79g3</u>
- 644
- 645 **Supplementary materials**
- 646 Tables S1-S3.
- 647 Fig. S1-S2.

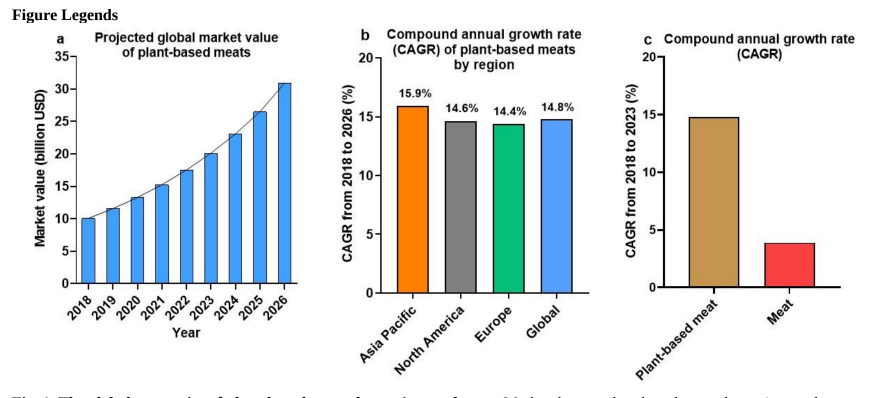
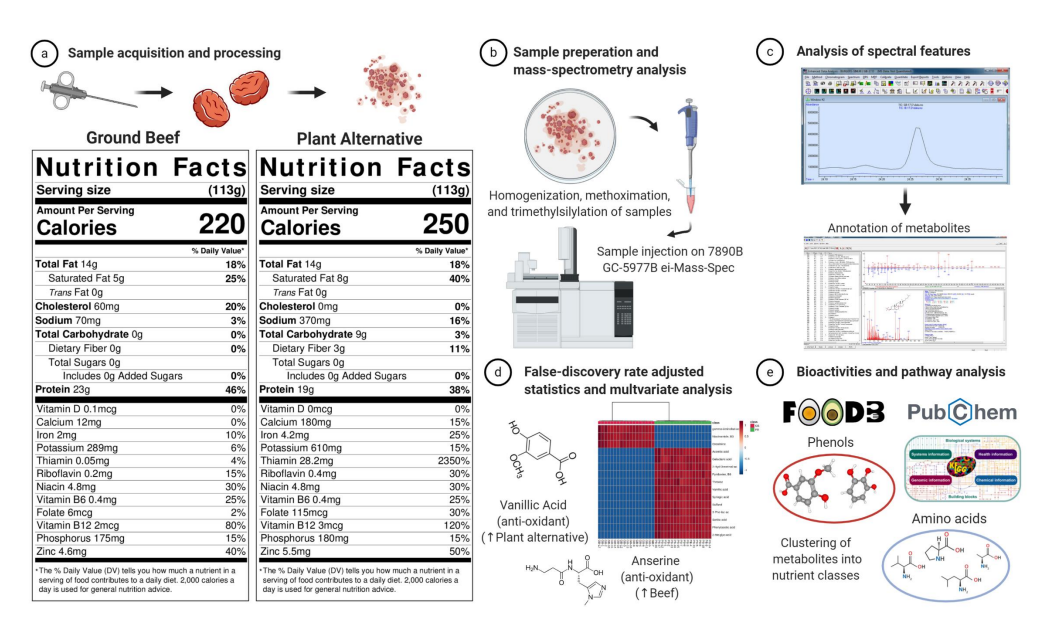


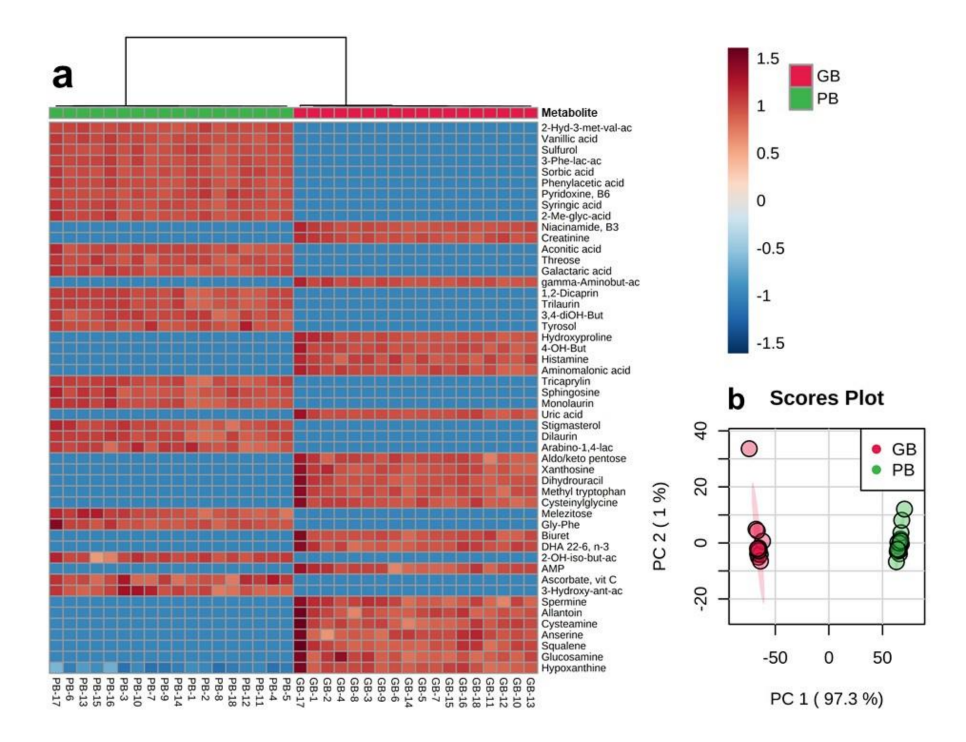
Fig. 1. The global economics of plant-based meat alternatives and meat. Market data on plant-based meat alternatives and meat were obtained from⁵. (**A**) The projected global market value of plant-based meats from 2018 to 2026 in Billion US Dollars. (**B**) The compound annual growth rate (CAGR) of the plant-based meat sector globally and by region. Amongst these regions, the largest growth is expected in the Asia Pacific. (**C**) The relative growth of the global plant-based meat sector (+14.8%) is expected to exceed the relative growth global animal meat market (+3.9%). Despite growth in absolute terms, the value share of the global animal meat

- 653 sector as a percentage of the overall food industry will remain more or less similar during 2018-2023⁵. This trend is due to a growing
- 654 preference among consumers for plant-based diets, which is motivated by concerns for human and environmental health⁵.



655	Fig 2. Schematic description of sample preparation and metabolomics analysis. (a) Nutrition Facts panels of grass-fed ground
656	beef and a market-leading plant-based meat alternative. Protein and fat content of the grass-fed ground beef was determined by
657	proximate analysis (Microbac Laboratories, Warrendale, PA), while the content of other nutrients in grass-fed beef were adapted from
658	US Department of Agriculture databases ⁶⁹ . Nutrient composition of the plant-based meat alternative was determined from its Nutrition
659	Facts panel. Eighteen burger patties of each product were cooked until an internal temperature of 71 °C, sampled using a bioptome,
660	and immediately frozen in liquid nitrogen (LN ₂) prior to further analysis. (b) Frozen samples were homogenized in 50% aqueous
661	acetonitrile containing 0.3% formic acid. Dried extracts were methoximated and trimethylsilylated, and untargeted metabolomic
662	analysis was conducted via gas chromatography/electron-ionization mass spectrometry (GC/ei-MS) on a 7890B GC-5977B ei-MS
663	(Agilent Technologies, Santa Clara, CA) in the Metabolomics Laboratory of the Duke Molecular Physiology Institute. (c) Raw
664	spectral data from Agilent's MassHunter software environment were imported into the freeware—Automatic Mass Spectral
665	Deconvolution and Identification Software or AMDIS. Peak annotation of metabolites was based primarily on our own RT-locked
666	spectral library of metabolites (2059 spectra from 1174 unique compounds). (d) To determine differences in abundance of metabolites
667	between beef and soy-based meat alternative, log-transformed metabolites were tested using the Wilcoxon rank sum test with
668	Benjamini-Hochberg adjusted <i>P</i> -values at 5% (False Discovery Rate; FDR < 0.05). (e) Bioactivities and potential health effects of
669	annotated metabolites were explored by entering metabolites in FooDB (https://foodb.ca/) and/or PubChem
670	(https://pubchem.ncbi.nlm.nih.gov/) databases, while metabolic pathway identification of individual metabolites was performed using
671	the Kyoto Encyclopedia of Genes and Genomes (KEGG) (https://www.genome.jp/). To further inform discussions of metabolomics

- 672 findings, metabolites were clustered according to structural similarly ChemRICH software procedures
- 673 (<u>http://chemrich.fiehnlab.ucdavis.edu/</u>). For further detail on these analyses see Methods section.



674	Fig. 3. Metabolomics revealed distinct differences in nutritional profiles between grass-fed ground beef (GB) and the plant-
675	based meat alternative (PB). (a) Heatmap of the top 50 metabolites, ranked by False Discovery Rate (FDR) adjusted P-values
676	(lowest to highest), that were significantly different (FDR < 0.05) between beef and the plant-based meat alternative. Red (intensity
677	ranges from 0 to 1.5) means higher abundance of the corresponding metabolite, whereas blue means lower abundance (intensity
678	ranges from -0 to -1.5). The numbers below the heatmap represent individual samples (GB-1 to 18 and PB-1 to 18 respectively; $n =$
679	18 for each group). Metabolites in beef and the plant-based meat were compared by the Wilcoxon rank sum test with Benjamini-
680	Hochberg adjusted <i>P</i> -values at 5% (FDR < 0.05). (b) Principal Component Analysis (PCA) analysis of beef and plant-based meat
681	revealed a distinct difference in nutritional composition between the grass-fed ground beef and the plant-based meat, with 97.3% of
682	the variance explained within the first principal component (PC1)—which illustrates the large nutritional differences that exist
683	between beef and the plant-based meat. The 95% confidence interval of the groups is depicted in each color. Red and green colors
684	above the heatmap (a) and the PCA plot (b) represent the ground beef and the plant-based meat, respectively. A full list of potential
685	bioactivities and health effects of each individual metabolite is reported in Table S1.

Table 1. Metabolites clustered into nutrient classes according to structural similarity using ChemRICH software procedures. Arrow

Nutrient Class	Class	No. different	↑ Plant	↑ Beef	FDR	Key Compound	Metabolic pathway, bioactivities/potential health effects
	size	plant vs beef	based				
Amino acids	19	18	12	6	<.001	Glutamine (†Plant)	Protein metabolism, neurotransmitter, anti-sickling, anti-ulcer
Non-protein amino acids	14	10	5	6	<.001	Creatinine (†Beef)	Energy metabolism, antioxidant, neuroprotective, ergogenic
Saccharides	13	12	8	4	<.001	Keto pentose-5-phos (<i>†</i> Beef)	Energy metabolism, flavor
Saturated fatty acids	11	9	3	6	<.001	Pentadecanoic acid (†Beef)	Odd-chain fatty acid biosynthesis, anti-bacterial, anti-oxidant,
Dicarboxylic acids	10	10	3	7	<.001	Aminomalonic acid (<i>†</i> Beef)	Glycine metabolism, unknown
Phenols	10	10	7	3	<.001	Vanillic acid (↑Plant)	Plant/microbial metabolism, anti-bacterial, anti-inflammatory
Dipeptides	8	6	2	4	<.001	Anserine (†Beef)	Carnosine metabolism, antioxidant
Purines	7	7	3	4	<.001	Uric acid (†Beef)	Microbrial/purine metabolism, unknown
Sugar alcohols	7	6	4	2	<.001	Myoinositol (Beef)	Biosynthesis, cholesterolytic, liver-protective, neuro-protective
Hydroxybuyrates	6	6	4	2	<.001	4-Hydroxybutyric acid (†Beef)	Biosynthesis, neurotransmitter, neuroprotective
Vitamins	5	5	3	2	<.001	Vitamin C (Plant)	Biosynthesis, anti-oxidant, liver-protective, kidney-protective
Glycerides	5	4	4	0	<.001	Monolaurin (†Plant)	Lipid metabolism, anti-microbial, anti-inflammatory
Pentoses	4	4	2	2	<.001	Arabinose/aldopentose (†Beef)	Energy metabolism, antioxidant, flavor
Sugar acids	4	4	3	1	<.001	Glyceric acid (†Beef)	Biosynthesis, cholesterolytic, diuretic, kidney-protective
Unsaturated fatty acids	4	4	2	2	<.001	Sorbic Acid (↑Plant)	Fatty acid biosynthesis, preservative
Amino alcohols	4	4	3	1	<.001	Phosphoethanolamine (<i>†</i> Beef)	Sphingolipid metabolism, neurotransmitter
Pyrimidines	4	3	1	2	.001	Dihydrouracil (↑Beef)	Pyrimidine metabolism, neuro-protective
Amines	4	3	0	3	.001	Cysteamine (†Beef)	Taurine metabolism, antioxidant, neuroprotective
Phytosterols	3	3	3	0	.003	Stigmasterol (†Plant)	Biosynthesis, anti-inflammatory, antioxidant, cancer-protective
Tocopherols	3	3	3	0	.003	γ-Tocopherol (↑Plant)	Biosynthesis, antioxidant, cardio-protective, cancer-protective
Biogenic polyamines	3	3	2	1	.003	Spermidine (Plant)	Glutathione metabolism, antioxidant
Polyunsaturated fatty acids	3	2	0	2	.008	DHA, 22-6, ω-3 (↑Beef)	Essential fatty acid, neuroprotective, cardio-protective
Pyridines	3	2	0	2	.017	3-Hydroxypyridine (†Beef)	Maillard reaction end-product, flavor
Fatty acid esters	3	1	1	0	1.00	1,2-Dicaprin (†Plant)	Energy metabolism, biosynthesis

([†]) indicates higher abundance for a particular nutrient class or nutrient.

DHA, docosahexaenoic acid; phos, phosphate.