

1    **Title**

2    Functional Anabolic Network Analysis of Human-associated *Lactobacillus* Strains

3    **Authors**

4    Thomas J. Moutinho Jr.<sup>a</sup>, Benjamin C. Neubert<sup>a</sup>, Matthew L. Jenior<sup>a</sup>, Maureen A. Carey<sup>a,b</sup>, Gregory L.  
5    Medlock<sup>a</sup>, Glynis L. Kolling<sup>a</sup>, Jason A. Papin<sup>a</sup>#

6    <sup>a</sup> Department of Biomedical Engineering, University of Virginia, Charlottesville, Virginia, USA

7    <sup>b</sup> Division of Infectious Disease and International Health, Department of Medicine, University of Virginia,  
8    Charlottesville, Virginia, USA

9    Running Head: Anabolic Network Analysis of Lactobacilli

10

11    #Address correspondence to Jason A. Papin, papin@virginia.edu

12    **Abstract**

13    Members of the *Lactobacillus* genus are frequently utilized in the probiotic industry with many species  
14    conferring demonstrated health benefits; however, these effects are largely strain-dependent. We  
15    designed a method called PROTEAN (Probabilistic Reconstruction Of constituent Anabolic Networks) to  
16    computationally analyze the genomic annotations and predicted metabolic production capabilities of  
17    144 strains across 16 species of *Lactobacillus* isolated from human intestinal, oral, and vaginal body  
18    sites. Using PROTEAN we conducted a genome-scale metabolic network comparison between strains,  
19    revealing that metabolic capabilities differ by isolation site. Notably, PROTEAN does not require a well-  
20    curated genome-scale metabolic network reconstruction to provide biological insights. We found that  
21    predicted metabolic capabilities of lactobacilli isolated from the vaginal microbiota cluster separately  
22    from intestinal and oral isolates, and we also uncovered an overlap in the predicted metabolic  
23    production capabilities of intestinal and oral isolates. Using machine learning, we determined the most  
24    informative metabolic products driving the difference between predicted metabolic capabilities of  
25    intestinal, oral, and vaginal isolates. Notably, intestinal and oral isolates were predicted to have a higher  
26    likelihood of producing D-alanine, D/L-serine, and L-proline, while the vaginal isolates were  
27    distinguished by a higher predicted likelihood of producing L-arginine, citrulline, and D/L-lactate. We  
28    found the distinguishing products to be consistent with published experimental literature. This study  
29    showcases a systematic technique, PROTEAN, for comparing the predicted functional metabolic output  
30    of microbes using genome-scale metabolic network analysis and computational modeling and provides  
31    unique insight into human-associated *Lactobacillus* biology.

32    **Importance**

33    The *Lactobacillus* genus has been shown to be important for human health. Lactobacilli have been  
34    isolated from human intestinal, oral, and vaginal sites. Members of the genus contribute significantly to  
35    the maintenance of vaginal health by providing colonization resistance to invading pathogens. A wide  
36    variety of clinical studies have indicated that *Lactobacillus*-based probiotics confer health benefits for

37 several gut- and immune-associated diseases. Microbes interact with the human body in several ways,  
38 including the production of metabolites that influence physiology or other surrounding microbes. We  
39 have conducted a strain-level genome-scale metabolic network reconstruction analysis of human-  
40 associated *Lactobacillus* strains, revealing that predicted metabolic capabilities differ when comparing  
41 intestinal/oral isolate to vaginal isolates. The technique we present here allows for direct interpretation  
42 of discriminating features between the experimental groups.

### 43 Introduction

44 *Lactobacillus* is a diverse genus of bacteria with many member strains associated with the human body.  
45 Lactobacilli are Gram-positive, lactic acid-producing bacteria typically with a low GC content (1,2). They  
46 are known for their production of lactic acid, being facultative anaerobes, and are capable of being  
47 metabolically active in a large variety of conditions (3). There is evidence that human-associated  
48 lactobacilli colonize mucosal surfaces of the intestinal tract (4), vagina (5–12), and oral cavity (13,14).  
49 While strains of *Lactobacillus* have been isolated from all three of these body sites, it remains unknown  
50 which are permanent members of the resident microbiota (autochthonous) opposed to transient  
51 members (allochthonous). Transient intestinal lactobacilli are either resident members of the oral  
52 microbiota or have been ingested, most commonly from unpasteurized fermented foods (4,15).

53 Lactobacilli have been used for a broad range of applications primarily associated with human intestinal  
54 probiotics and industrial production of useful metabolites. *Lactobacillus*-based probiotics have been  
55 shown to confer health benefits in clinical studies for a variety of conditions including prevention of  
56 antibiotic associated diarrhea (16), *Clostridium difficile*-associated diarrhea (17), constipation (18),  
57 irritable bowel syndrome (19), and eczema/atopic dermatitis (20). Probiotics are controversial, likely due  
58 to claims made by currently marketed probiotics that lack FDA approval for the treatment of specific  
59 diseases (21,22). The primary benefits associated with lactobacilli-based probiotics may be a function of  
60 their presence in the gut, production of metabolites, and modulation of the immune system (23,24).  
61 Metabolism plays a key role in all three of these general mechanisms; therefore, a better understanding  
62 of their metabolic capabilities will help to elucidate the mechanisms contributing to probiotic effects  
63 (25).

64 In recent years, there has been an explosion of genomic and metagenomic sequencing of human-  
65 associated microbiota, which provides a unique opportunity to apply genome-scale metabolic network  
66 reconstructions (GENREs) to enhance our current understanding of human-associated lactobacilli  
67 metabolism utilizing *in silico* techniques (25). Systems biology has the potential to advance design,  
68 selection, and delivery of *Lactobacillus*-based probiotics (26,27). GENREs are a powerful computational  
69 tool for mathematically modeling the metabolic processes within a cell at a systems-level, including all  
70 known metabolic reactions, metabolites, and metabolic genes in an organism (28). GENREs are created  
71 by referencing an annotated genome against biochemical databases, then integrating experimental data  
72 when available (29). There are several examples of *Lactobacillus*-specific comparative genomics studies  
73 (30–35); however, GENREs allow for a more functional perspective than genomics data alone because of  
74 the quantitative accounting for interactions between components in the network (25,36). Simulations  
75 with GENREs can accurately predict microbial growth yields and the metabolic pathways utilized for the  
76 production of metabolites during exponential growth of a microbe (37). A variety of analytical  
77 approaches can be applied to interrogate emergent properties of a GENRE. Flux Balance Analysis (FBA)  
78 and related methods have proven highly successful in the analysis of metabolic networks (38). FBA is a

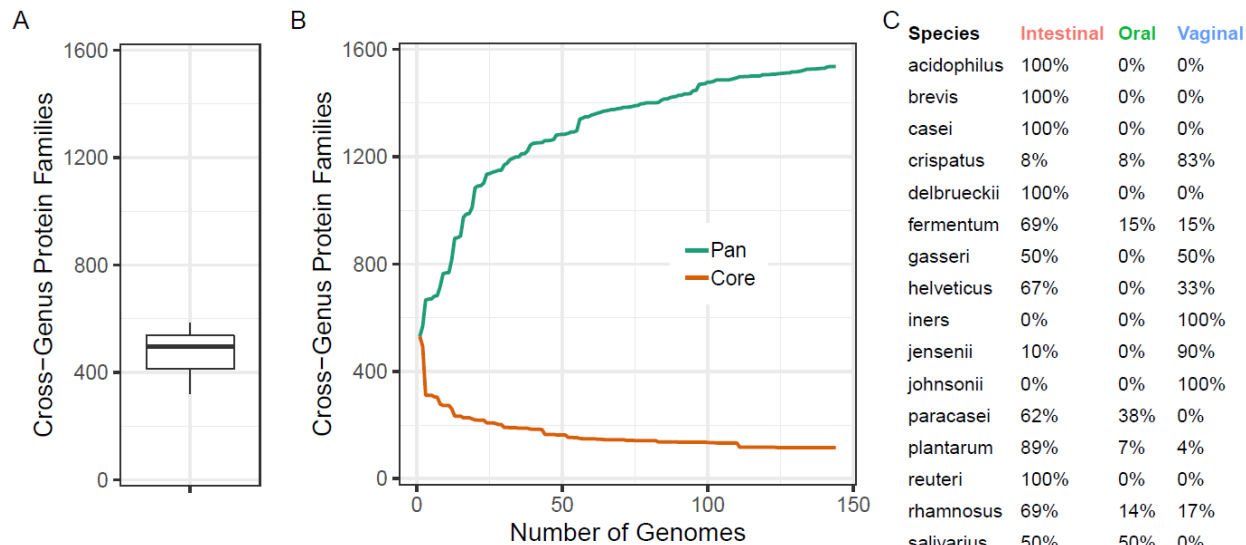
79 mathematical technique for analyzing the flow of metabolites through a GENRE; it can be used to  
80 identify a set of reaction fluxes that maximize growth in a specified media condition among other  
81 applications (28,39,40). Metabolic network reconstructions and FBA provide a mechanistic look into  
82 cellular metabolism and are increasingly used to study biochemical processes of single bacterial species  
83 as well as communities of organisms (41).

84 GENREs enable the computational prediction of metabolic capabilities of microbes, both catabolic and  
85 anabolic. Additionally, GENREs are capable of contextualizing large 'omic datasets (i.e. genomics,  
86 transcriptomics, and metabolomics) with known biochemistry and biological network architectures for  
87 improved understanding of the experimental data (42). An important recent finding demonstrated that  
88 metabolomics data alone can be used to differentiate between bacterial cultures at the strain level (43).  
89 We developed a computational method using GENREs to predict the metabolic products that a strain is  
90 likely able to produce. We used predicted production capabilities to then differentiate between  
91 different human-associated *Lactobacillus* strains. Just as metabolomics data can be used to differentiate  
92 bacterial strains, predicted production capabilities can be used for the same comparisons. We assessed  
93 the metabolic potential across a broad set of *Lactobacillus* species, consisting of 144 strains, which have  
94 all been isolated from three human-related body sites: intestinal, oral, and vaginal. We found that  
95 intestinal and oral isolates have a great deal of overlap in their metabolic functionality, while vaginal  
96 isolates have more unique metabolic production capabilities. These analyses can facilitate additional  
97 experimental interrogation of this important genus of bacteria.

## 98 Results and Discussion

### 99 Annotated metabolic genes associated with known metabolic functions are sufficiently represented 100 among human-associated lactobacilli

101 In this study we predict the metabolic production capabilities of 144 lactobacilli strains. We utilized the  
102 PATRIC Cross-Genus Protein Families (PGfams) (4) for an initial genomic analysis. PGfams are  
103 comparable clusters of proteins that likely have similar functions. These clusters are intended to be used  
104 for cross-genus comparison due to their slightly relaxed clustering criteria. However, PGfams allow for  
105 the comparison of the large number of strains analyzed in this study. Lactobacilli consist of a broad  
106 range of species and thus using the PGfams was appropriate for an initial genomic comparison in this  
107 study. We first filtered the PGfams to only include metabolic gene families associated with known  
108 metabolic functions (see Methods). The distribution of total metabolic PGfams associated with each  
109 genome ranges from 340 to 580 and has a median value of 515 (Figure 1A). Across these 144 strains we  
110 found that they share 116 core metabolic PGfams, spanning a variety of cellular functions including, but  
111 not limited to, carbohydrate, nucleotide, and amino acid metabolism (Table S1). The pan set of  
112 metabolic PGfams, which represents the total set of unique PGfams, expanded to over 1500 after  
113 considering all strains utilized within this study (Figure 1B). The *Lactobacillus* strains we studied  
114 consisted of 16 species and were isolated from intestinal, oral, and vaginal human body sites (Figure 1C).



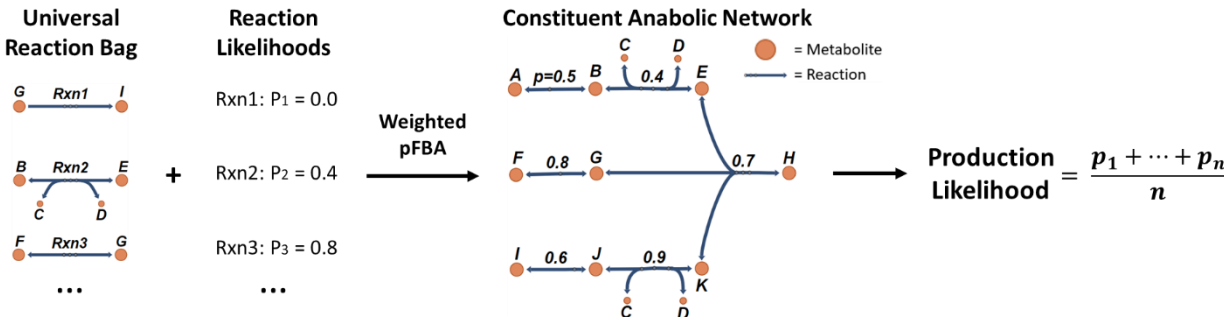
115

116 **Figure 1: Known metabolic annotations are extensively sampled across the 16 *Lactobacillus* species included in**  
117 **this study.** The genomic features used for this analysis are PATRIC Cross-Genera Protein families (PGfams), a  
118 standard set of features across the PATRIC Database (4). (A) The number of metabolic PGfams for each  
119 genome are shown here, with the median value indicated by the middle line in the boxplot. (B) For the 144 strains  
120 from 16 species of *Lactobacillus*, we found that there are 116 protein families in the core set of metabolic PGfams,  
121 while the pan set of PGfams expands to over 1500 families. The nearly plateau shape of the curve for the pan set  
122 of PGfams curve indicates that this sampling represents a large portion of the genetic diversity among the 16  
123 species included in the study. (C) This table shows the complete list of species used in this study and indicates the  
124 percentage of strains that were isolated from each human body site. Each strain in this study is a member from  
125 one of the 16 species and isolated from one of three human-associated body sites; intestinal, oral, or vaginal (Table  
126 S2).

## 127 Probabilistic Reconstruction Of constituent Anabolic Networks (PROTEAN)

128 We developed PROTEAN to predict the metabolic production capabilities of microbes based on genomic  
129 data alone. PROTEAN generates constituent metabolic production networks with maximum parsimony  
130 and probability to predict the production of a given metabolite with a defined set of input metabolites.  
131 PROTEAN is a combination of well-validated methods, including Parsimonious Enzyme Usage Flux  
132 Balance Analysis (pFBA) (37), likelihood-based gap filling (44), fastGapFill (45), and CarveMe (46). The  
133 algorithm uses the ModelSEED biochemical reaction database, a large set of known metabolic reactions,  
134 for constituent network generation (47). First, reaction likelihoods are calculated for each reaction in the  
135 ModelSEED database using Probannopy (48) (Figure 2). Reaction likelihoods correspond to the  
136 probability that a given reaction is catalyzed by an enzyme that is encoded for by the genome. We  
137 modified pFBA to utilize reaction likelihoods for weighted minimization of flux through each reaction,  
138 while still maintaining near-optimal flux through the objective function. Standard pFBA assumes that  
139 metabolism is optimized to minimize enzymatic turnover and thus the method is driven by a  
140 minimization of the total flux through the metabolic network (37). Weighted pFBA allows for the  
141 reconstruction of constituent anabolic networks while accounting for maximum genomic probability and  
142 resource parsimony (see Methods). The constituent anabolic networks output by PROTEAN consist of  
143 flux-carrying reactions required for the production of a certain metabolite with preferential flux through  
144 reactions that have higher reaction likelihoods. A constituent network represents a theoretically optimal

145 biosynthetic network while accounting for the greatest genomic evidence for production of a given  
146 metabolite in a set media condition (Table S4). We represent the information from each constituent  
147 network using a single summary metric referred to as the Production Likelihood by calculating the  
148 average of all likelihoods of reactions that carry flux. The average of all reaction likelihoods in a  
149 metabolic pathway has been previously shown to be a valuable metric for making comparisons between  
150 networks (44).

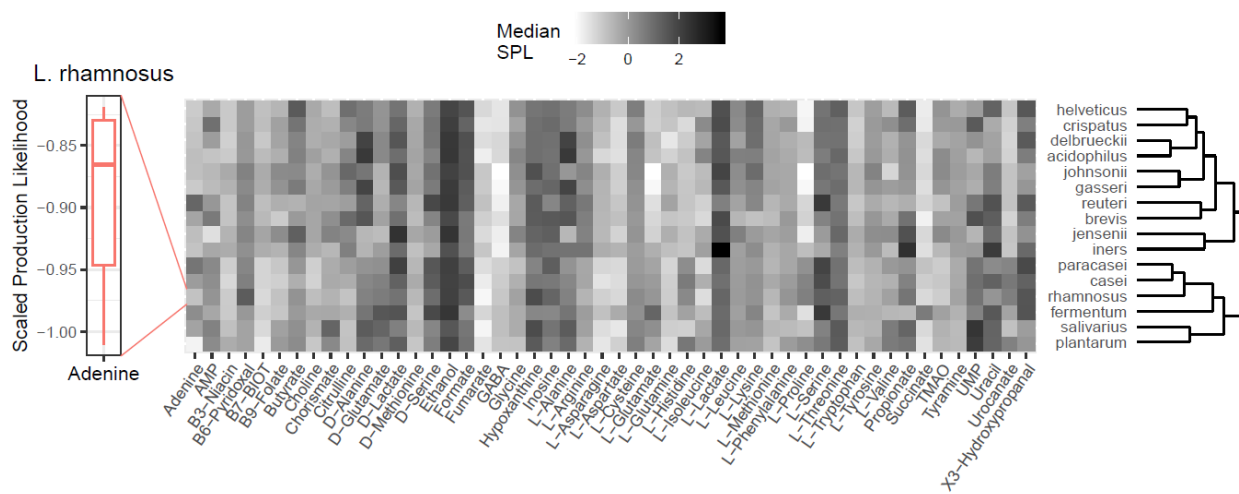


151  
152 **Figure 2: PROTEAN is an approach for quantifying the likelihood that a given metabolic network, derived**  
153 **exclusively from genomic evidence, is capable of synthesizing a particular metabolite.** A modified version of  
154 Parsimonious Enzyme Usage FBA (weighted pFBA) was performed on a standardized set of reactions to generate  
155 constituent anabolic networks for each genome. Reaction likelihoods were used to weight the minimization of flux  
156 through each reaction in the network. Therefore, reactions with a greater likelihood were more likely to be  
157 included in the resulting constituent anabolic network. Each constituent network has a set of input metabolites  
158 representing the media condition (Table S4) and a demand reaction for a certain metabolic product. The resulting  
159 constituent network is the set of reactions that requires flux to produce the metabolic product in the given media  
160 condition. The production likelihood metric is an average of all the reaction likelihoods associated with the  
161 reactions included in the constituent network. This metric is used as a summary statistic that allows for the  
162 comparison of constituent networks across different metabolic products and strains, where a higher production  
163 likelihood corresponds with greater genetic evidence for that particular constituent anabolic network.

164 **The Scaled Production Likelihood metric facilitates comparison of anabolic capabilities between**  
165 **species and strains**

166 Predicted constituent anabolic networks were generated for a set of 50 biologically-relevant metabolic  
167 products for each of the 144 *Lactobacillus* strains. The 50 metabolites were selected based on known  
168 *Lactobacillus* biology (see Methods). For each metabolic product, we generated a constituent anabolic  
169 network (Table S3) across all strains. For each genome we scaled the Production Likelihoods metric by  
170 calculating the corresponding z-score. The standard deviation for the z-score calculation was across all  
171 metabolic products for each strain. This metric allows for a relative comparison of production  
172 capabilities across strains that does not rely on well-curated metabolic network reconstructions. The  
173 resulting Scaled Production Likelihood (SPL) is a metric indicating likelihood that a genome encodes for  
174 the cellular machinery required to produce a metabolite, given a specific media condition, relative to all  
175 of the other SPLs for the metabolic products per strain. For visualization, these data were grouped by  
176 species and summarized using the median of the SPLs across all of the strains within each species (Figure  
177 3).

178



179

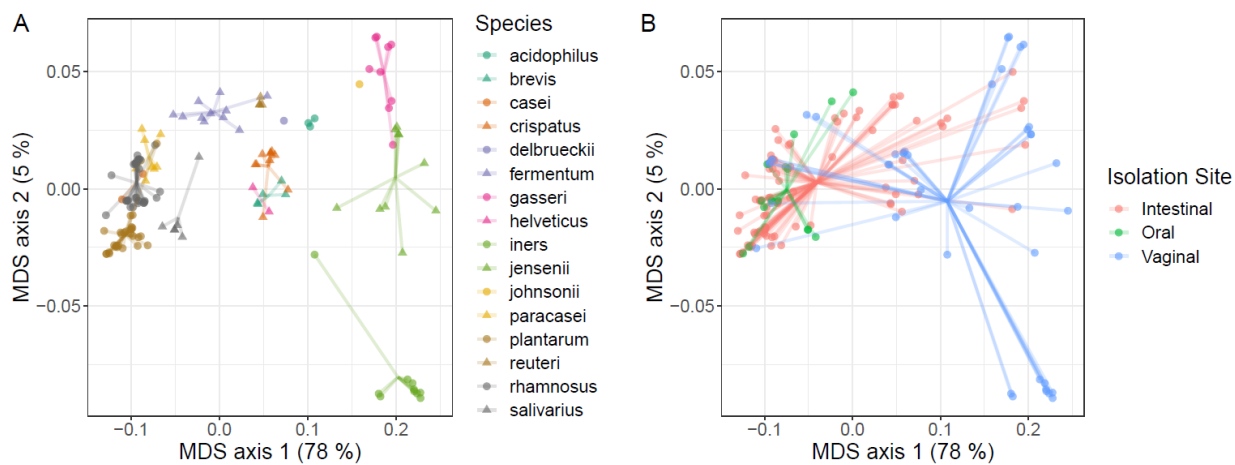
180 **Figure 3: Predicted metabolic production capabilities with the Scaled Production Likelihood (SPL) metric align**  
181 **poorly with phylogeny.** There is a single production likelihood for each genome associated with each metabolite. A  
182 median SPL can be calculated for a species that allows for more general comparisons across species, illustrated  
183 here by the distribution for one species (*L. rhamnosus*) and one metabolite (adenine). There are 50 metabolites  
184 used as features to allow for the comparison of predicted production capabilities across the lactobacilli analyzed.

185 The strains were grouped by species and clustered based on median SPLs. We found that across the 16  
186 species, D- and L-lactate both have high median SPLs, as we would expect with lactobacilli. Additionally,  
187 fumarate and GABA have particularly low SPLs across all species. We were able to find several  
188 publications indicating GABA can be produced by select lactobacilli in specific environments (49,50).  
189 However, we were unable to find publications discussing the production of fumarate by lactobacilli.  
190 Additionally, we found that the dendrogram from clustering based on predicted metabolic production  
191 capabilities does not qualitatively align well with published phylogenetic trees generated using the 16S  
192 rRNA gene (34). The misalignment to established phylogenetic trees indicates that phylogeny is a poor  
193 indicator of metabolic production capabilities. It is likely that evolution of metabolic production  
194 capabilities is driven independently from classical genes used for phylogenetic comparisons, such as the  
195 16S rRNA gene. Therefore, we need more precise computational tools to better understand the  
196 phenotypic differences between microbial species when interrogating metabolism. Perhaps  
197 phylogenetic analysis would be augmented with the consideration of metabolic genes in addition to the  
198 16S rRNA gene.

199 **Intestinal and oral *Lactobacillus* strains have different metabolic capabilities compared to vaginal**  
200 **strains**

201 We performed principle coordinate analysis (PCoA) on the SPLs for each species and determined that  
202 the *Lactobacillus* strains cluster significantly by both species (Figure 4A) and isolation site (Figure 4B)  
203 (PERMANOVA;  $P < 0.001$ ). The vaginal isolates differ from both the oral and gut cluster (Figure 4B).  
204 Substantial overlap was found between oral and gut isolates, specifically within *L. gasseri*, *L. rhamnosus*,  
205 and *L. salivarius*, likely due to the consistent transmission of orally colonized microbes to the intestines  
206 (15). It has been hypothesized that many of the lactobacilli isolated from the gut are actually transient  
207 strains that are colonized in the oral cavity (51). Our data supports this hypothesis by showing that oral

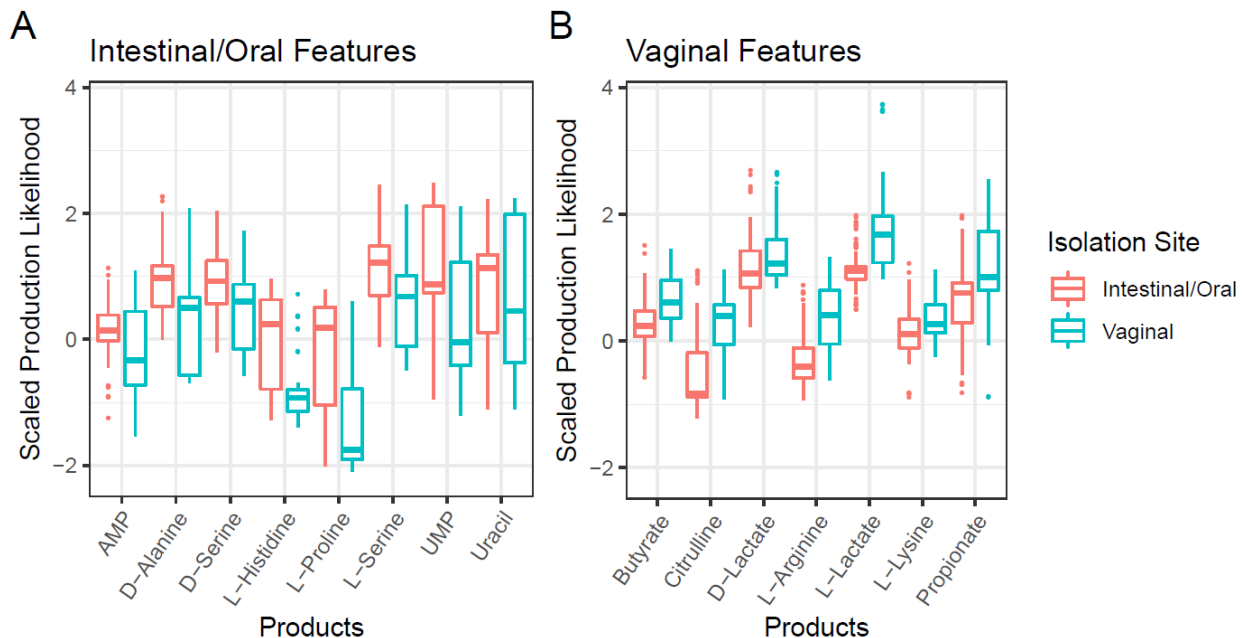
208 isolates are metabolically similar to a portion of the intestinal isolates. However, there are lactobacilli,  
209 such as *L. reuteri*, which likely colonize the human intestines (52). Five of the 16 species in this study are  
210 only represented by strains isolated from the intestines; although this result is influenced by sampling  
211 bias in the PATRIC Database, it provides support that our data contains species that are only found in  
212 the intestines. The vaginal isolates cluster separately from the intestinal/oral isolates along the primary  
213 coordinate that accounts for 78% of the variation in these data. The vaginal microbiota is frequently  
214 dominated by several *Lactobacillus* species, such as *L. iners*, *L. crispatus*, and *L. jensenii* (53–55). This  
215 separation of vaginal isolates from intestinal/oral isolates indicates that these two main clusters have  
216 differences in their metabolic production capabilities. This result is to be expected because the  
217 intestinal/oral nutrient environment is drastically different from the vaginal environment and the  
218 dominant species appear to have metabolic capabilities that reflect this difference.



219  
220 **Figure 4: The Scaled Production Likelihood metric distinguishes metabolic functionality among species.** (A) We  
221 found that *Lactobacillus* strains cluster significantly by species (PERMANOVA;  $P < 0.001$ ). (B) Additionally, they  
222 cluster significantly by isolation site (PERMANOVA;  $P < 0.001$ ). Both plots are PCoA using the Bray-Curtis distance  
223 metric of the SPLs for each isolate. Points in both panels are identical, but displayed with different color schemes.

224 In addition to distinguishing isolates by body site, the SPL metric is capable of defining collections of  
225 functional components that drive differences between groups. Using standard genomic analyses,  
226 differences between groups are typically defined by the differential gene content. Genes are intrinsically  
227 part of a larger network of metabolism where absence of specific functionality related to a gene may be  
228 compensated for within the system. Since our approach is based on Production Likelihoods of specific  
229 metabolites, it functions within a more complex metabolic framework compared to the analysis of  
230 genomic data without the network context. Using machine learning, we were able to identify the set of  
231 metabolites for which each group of strains is more likely to encode the cellular machinery required for  
232 production. We conducted a machine learning feature selection to determine the metabolites that are  
233 most likely to be produced by each group of strains, intestinal/oral strains and vaginal strains. We  
234 grouped the intestinal and oral strains together due to their inherent similarity (Figure 4B) and the  
235 observed transmission of oral strains to the intestines (15,51). We generated two separate area under  
236 the curve random forest (AUCRF) models to determine the metabolites that were more likely to be  
237 produced by each of the groups. Two models were necessary to enrich for the most discriminatory  
238 metabolites that were more likely to be produced in each of the groups, rather than simply identifying  
239 the metabolites that best classify the samples based on isolation site regardless of being more or less

240 likely to be produced (See methods). The first model was generated to select the metabolites that are  
241 most likely to be produced by the intestinal and oral isolates compared to the vaginal isolates, while  
242 maximally discriminating the groups. The eight metabolites selected accurately classify greater than 90%  
243 of isolates to the correct group (Figure 5A). The second model was generated to select the metabolites  
244 that are most likely to be produced by the vaginal isolates compared to the intestinal and oral isolates,  
245 while maximally discriminating the groups. The seven metabolites selected accurately classify greater  
246 than 90% of the isolates to the correct group (Figure 5B).



247

248 **Figure 5: Machine learning of the SPL scores identifies metabolites that discriminate *Lactobacillus* strains.**  
249 Machine learning feature selection identified the metabolites that are both most likely to be produced by each  
250 group and capable of classifying the strains into two groups, intestinal/oral and vaginal, with greater than 90%  
251 accuracy. (A) There are eight metabolites that are more likely to be produced by the intestinal/oral isolates  
252 compared to the vaginal isolates. (B) There are seven metabolites that are more likely to be produced by the  
253 vaginal isolates compared to intestinal/oral isolates. Both models are more than 90% accurate in predicting the  
254 membership to which the given isolate belongs using the SPLs of the metabolites listed.

255 Using SPLs as an input for AUCRF feature selection, we identified the metabolites that are most likely to  
256 be produced by the strains associated with the two isolate groups, intestinal/oral and vaginal. The  
257 selected metabolite products may contribute to how the strains interact with the mucosal tissues in  
258 each site. We hypothesize that these metabolites are related to key phenotypic differences between the  
259 two isolate groups. Four of the selected metabolites that are likely produced by intestinal/oral strains,  
260 D-alanine, D/L-serine, and L-proline (Figure 5A), have all been previously identified to have an impact on  
261 the human intestinal epithelium (23,24,56–58). Additionally, four of the selected metabolites that are  
262 likely produced by vaginal strains, L-arginine, citrulline, and D/L-lactate (Figure 5B), have been previously  
263 identified to have an impact on the human vaginal microbiome (59–62). The metabolites for which we  
264 have not found existing experimental evidence for are likely worth focusing on in future experimental  
265 studies.

266 For intestine-associated lactobacilli in this study, there is a connection between intestinal immune  
267 system regulation and D-alanine rich lipotechoic acid, a glycolipid expressed by some lactobacilli, such as  
268 *L. plantarum* (23,24). D-alanine rich lipotechoic acid, produced by lactobacilli, has been shown to down-  
269 regulate local colonic inflammation in a murine colitis model (23,24). With PROTEAN we identified that  
270 intestinal lactobacilli were more likely to produce D-alanine (Figure 5A). It is possible that a positive  
271 interaction with the intestinal host immune system would result in an evolutionary advantage by  
272 reducing local immune response. Additionally, serine rich serine-threonine peptides have been shown to  
273 have a similar regulatory effect on intestinal dendritic cells (56,57). These peptides expressed by *L.*  
274 *plantarum* are resistant to intestinal proteolysis and appear to be present in the colon of most healthy  
275 individuals (56,57). Similar to D-alanine, the production of D/L-serine would require a robust  
276 biosynthesis pathway present in those strains.

277 A final gut-related connection involves the biosynthesis of L-proline (Figure 5A). One of the primary  
278 stress responses in *L. acidophilus* to high osmotic pressure results in the accumulation of L-proline in the  
279 cell; there is little evidence that this response is a result of L-proline transport into the cell (58). These  
280 *Lactobacillus* strains are exposed to a large range of stressors in the gut, including suboptimal osmotic  
281 pressures. There is strong evidence that L-proline is used by *L. acidophilus* to tolerate suboptimal  
282 osmotic pressures and there is a lack of evidence for L-proline transporters. As such, the biosynthesis of  
283 L-proline may be advantageous for growth in the gut.

284 For the enriched metabolic products in vaginal isolates (Figure 5B), there is evidence for an  
285 arginine/ornithine antiporter and arginine deiminase in *L. fermentum* (59). These enzymes are part of  
286 the arginine deiminase pathway through which there is the production of citrulline which is exported  
287 from the cell and contributes to acid tolerance (59). It has also been demonstrated that treatment with  
288 probiotics containing arginine deiminase-positive lactobacilli can improve clinical symptoms of vaginosis  
289 in parallel with significant declines in polyamine (i.e. arginine, ornithine, and citrulline) levels in the  
290 vagina (60,61). The vaginal isolates in this study show enrichment for the cellular machinery required for  
291 the production of both citrulline and L-arginine (Figure 5B). The importance of lactate for the adequate  
292 maintenance of vaginal health in many individuals is known. The current hypothesis revolves around  
293 colonization resistance where vaginal lactobacilli establish an acidic environment by producing lactate  
294 (62). The acidic environment is generally inhospitable to invading pathogens as well as other microbes  
295 that are otherwise capable of residing in the vaginal environment (62). It has been shown that higher  
296 levels of D-lactate over L-lactate present in the vagina, produced by lactobacilli, further decrease the  
297 chance of infections in female patients (62). However, both isoforms of lactate remain important in  
298 maintaining vaginal health.

## 299 Conclusions

300 Microbial biosynthesis of metabolites has a broad range of applications, from bio-manufacturing to  
301 microbiome research (63). There is a wealth of well-curated and accessible knowledge stored in  
302 biochemical reaction databases such as ModelSEED (64). Genome-Scale Metabolic Network  
303 Reconstructions access this fundamental knowledge while accounting for systems-level interactions.  
304 This study represents one such application of GENREs that is a step toward predicting the metabolic  
305 production capabilities of understudied organisms. Experimental validation of the production  
306 capabilities predicted with PROTEAN will allow for conclusions to be made beyond the statement that a  
307 microbe is genetically likely to be able to produce a metabolite. Utilizing PROTEAN data, we found that

308 human-associated lactobacilli strains cluster significantly by species and isolation site. Additionally,  
309 many of the metabolic products that drive the clustering of strains by the isolation sites have known  
310 physiological function and importance in the respective isolation sites.

311 Future applications of PROTEAN could include optimal strain selection for bio-manufacturing of a certain  
312 compound, generating predicted metabolomics data for an organism to generate a prioritized list of  
313 conditions that would be most worthwhile to validate experimentally, and predicting the metabolites  
314 that are most likely to be produced in a microbiota. Microbes can have a wide range of physiological  
315 impacts on human health; these impacts are, in part, a result of the metabolites that are or are not  
316 produced by members of a microbiota. One of the core limitations of this study includes the lack of  
317 reaction likelihoods for some reactions in the universal reaction bag we used from ModelSEED. The  
318 number of reactions we could generate likelihoods for was limited by the Probannopy reaction  
319 template. However, this template can be expanded to continue to improve the utility of PROTEAN. With  
320 the inclusion of validation data, additional analyses will be possible, such as determining metabolic  
321 production pathways lacking proper annotation. By determining the reactions that are most likely  
322 required for biosynthesis of a known product, it would be possible to generate additional hypotheses for  
323 enzyme annotation experiments. PROTEAN is an algorithm with potential for a wide range of  
324 applications in the study and use of microbial metabolic networks.

## 325 Methods

### 326 Constituent Anabolic Network Generation (PROTEAN)

327 Probabilistic pFBA-based constituent anabolic network generation was accomplished using three Python  
328 packages, Cobrapy (65), Mackinac (66), and Probannopy (48). The complete ModelSEED universal  
329 reaction bag was downloaded from the github repository and filtered based on the annotation quality  
330 score, including all reactions with an 'OK' quality status or better (64). For each reaction in the  
331 ModelSEED universal reaction bag, we used Probannopy to generate a reaction likelihood based on the  
332 FASTA file for each genome obtained from the PATRIC database (4). The Cobrapy implementation of  
333 Parsimonious Enzyme Usage Flux Balance Analysis (pFBA) was altered to allow for each reaction's linear  
334 constraint to be set individually based on the reaction likelihood. The linear constraint for each reaction  
335 was set to one minus the reaction likelihood (a value between 0 and 1). There were reactions included in  
336 the universal reaction bag that were lacking from the Probannopy template model, therefore resulting  
337 in several gene-associated reactions lacking reaction likelihood scores. The reactions without likelihoods  
338 were left at a full minimization penalty (linear constraint value of 1). We chose to penalize the reactions  
339 without likelihoods to bias our results towards the construction of networks for which all reactions had  
340 evidence of presence. The linear constraints applied to each reaction based on likelihood acted as a  
341 weighting (inclusion penalty) for the minimization step in pFBA, resulting in the reactions with greater  
342 likelihood having a lower penalty for carrying flux; therefore, the reactions had a higher likelihood of  
343 being included in the constituent anabolic networks.

344 Using PROTEAN, we generated constituent anabolic networks by setting a certain input media condition  
345 (Table S4) and constraining flux through the single metabolite objective function (Table S3). We ran our  
346 likelihood-weighted pFBA flux minimization across the entire universal reaction bag and isolated the  
347 reactions that carried flux to get the desired product. The resulting networks consist of the direct  
348 reactions that would be part of a production pathway as might be shown in a typical biosynthesis  
349 pathway figure, while also accounting for all of the secondary and energy metabolites that are required

350 for the production of the metabolite in consideration. Additionally, this algorithm is optimizing for three  
351 core characteristics in the constituent networks: 1) minimum flux through the network (loosely, the  
352 minimum number of reactions), 2) maximum average reaction likelihood across the constituent  
353 network, and 3) output flux within 90% of the optimal yield of the metabolic product. We chose to allow  
354 flux through any reaction in the universal reaction bag during the generation of the constituent anabolic  
355 production pathways rather than simply pulling from a GENRE that was first gapfilled to allow  
356 production of biomass. Using the universal reaction bag instead of a gapfilled model was important  
357 because the biomass function is difficult to define for understudied organisms and unnecessary for our  
358 applications.

### 359 **Scaled Production Likelihood Metric**

360 We represent the information from each constituent network using a single summary metric for ease of  
361 comparison, simply named the Production Likelihood. This metric is the average of the reaction  
362 likelihoods included in the constituent network. The average reaction likelihood for a metabolic pathway  
363 has been previously used for making comparisons between networks (44). The Production Likelihoods  
364 for all 50 metabolites are scaled for each given genome by calculating the z-score to create the Scaled  
365 Production Likelihoods used for the majority of the analysis in this study. The z-score is calculated for  
366 each individual strain using the median and standard deviation for the production likelihoods across the  
367 50 metabolic products. The Scaled Production Likelihood allows for a ranked comparison of metabolic  
368 products across the genome set and corrects for annotation bias by essentially comparing the ranked z-  
369 score for each metabolic product.

### 370 **Supporting data for pathway generation**

371 The simulated media formulation was based on *in vitro* minimal media growth conditions for *L.*  
372 *plantarum* (Table S4) (67–69). The techniques used in this study do not assume that all species are  
373 capable of growth in the given media condition, therefore this media condition simply provides a  
374 standard reference for comparison. The product list was developed by identifying metabolites that have  
375 been shown to be produced by lactobacilli during *in vitro* growth experiments, in addition to other  
376 metabolites that have been shown to be related to human physiology (70–74).

### 377 **Machine learning feature selection**

378 Discriminating intestinal/oral and vaginal features were selected using area under the ROC curve  
379 random forest (AUCRF) using default parameters (75) (see Code). We generated two separate AUCRF  
380 models to determine the metabolites that were more likely to be produced by each of the groups,  
381 intestinal/oral and vaginal. Two models allowed us to enrich for likely products rather than simply  
382 selecting for the metabolites that provide the greatest discrimination between the groups but which  
383 may have poor likelihood scores. We conducted the enrichment for likely metabolic products for each  
384 model by reducing the feature set down to only metabolites that were more likely to be produced by  
385 the group of interest. Likely metabolic products were determined by comparing the median SPLs of each  
386 metabolite between the groups. Additionally, the feature sets were reduced to include only metabolites  
387 with a median value greater than zero for the group of interest. An AUCRF model was then generated to  
388 select the features that provided the greatest discrimination between the two groups.

### 389 **Statistical modeling and figure generation**

390 The principle coordinate analysis (PCoA) ordinations were created using the R vegan package (76),  
391 implemented with the Bray-Curtis dissimilarity metric. Statistical significance for comparing the PCoA  
392 clusters was determined using a PERMANOVA (R Adonis test). A variety of R packages were used for all  
393 figure generation (77–81).

#### 394 **Genome Quality and PATRIC Cross Genus Protein Family Data**

395 Genomes used in the study were filtered for quality before being included in the analysis. Strains with  
396 greater than 0.2% unknown nucleotide calls in the genome were eliminated. Low quality genome  
397 assemblies with greater than 300 contigs were removed. Non-human associated *Lactobacillus* strains  
398 from the PATRIC database were used to determine the GC content range for each species (82,83), and  
399 significant outliers (plus or minus two percent) were removed to control for sequencing bias (84,85).  
400 Only isolates from the three human-associated sites (oral, intestinal, and vaginal) were included in the  
401 final dataset.

402 The inclusion of metabolic PATRIC cross genus protein families was conducted by filtering the PGfams  
403 for each genome based on the existence of an associated known reaction and ProbannoP likelihood  
404 greater than 0. Pan and core metabolic PGfam sets were evaluated after the addition of all genomic  
405 features from each genome. The pan set of metabolic PGfams was defined as the total number of  
406 unique PGfams included in the data set after the above filtering steps. The core set of metabolic PGfams  
407 are those that existed within each genome included in this study.

#### 408 **Data and code availability**

409 Genome FASTA files and metadata were downloaded from the PATRIC Database (4). Python and R code  
410 is available at: [Github.com/Tjmoutinho/Lactobacillus](https://github.com/Tjmoutinho/Lactobacillus)

## 411 **References**

- 412 1. de Vos WM. Systems solutions by lactic acid bacteria: from paradigms to practice. *Microb Cell*  
413 *Factories*. 2011 Aug 30;10(1):S2.
- 414 2. de Vos WM, Hugenholtz J. Engineering metabolic highways in Lactococci and other lactic acid  
415 bacteria. *Trends Biotechnol*. 2004 Feb 1;22(2):72–9.
- 416 3. Ljungh Å, Wadström T. *Lactobacillus Molecular Biology: From Genomics to Probiotics*. Horizon  
417 Scientific Press; 2009. 217 p.
- 418 4. Wattam AR, Abraham D, Dalay O, Disz TL, Driscoll T, Gabbard JL, et al. PATRIC, the bacterial  
419 bioinformatics database and analysis resource. *Nucleic Acids Res*. 2014 Jan 1;42(D1):D581–91.
- 420 5. OHanlon DE. In vivo versus in vitro metabolomics profiling of vaginal lactobacilli for probiotic use.  
421 2013 Jun 4 [cited 2018 Sep 24]; Available from: <https://www.omicsonline.org/proceedings/in-vivo-versus-in-vitro-metabolomics-profiling-of-vaginal-lactobacilli-for-probiotic-use-785.html>
- 423 6. O’Hanlon DE, Moench TR, Cone RA. Vaginal pH and Microbicidal Lactic Acid When Lactobacilli  
424 Dominate the Microbiota. *PLoS ONE* [Internet]. 2013 Nov 6 [cited 2018 Sep 24];8(11). Available  
425 from: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3819307/>

- 426 7. Tachedjian G, Aldunate M, Bradshaw CS, Cone RA. The role of lactic acid production by probiotic  
427 Lactobacillus species in vaginal health. *Res Microbiol.* 2017 Nov 1;168(9):782–92.
- 428 8. Tachedjian G, O'Hanlon DE, Ravel J. The implausible "in vivo" role of hydrogen peroxide as an  
429 antimicrobial factor produced by vaginal microbiota. *Microbiome [Internet].* 2018 Feb 6 [cited 2018  
430 Sep 24];6. Available from: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5801833/>
- 431 9. Parolin C, Foschi C, Laghi L, Zhu C, Banzola N, Gaspari V, et al. Insights Into Vaginal Bacterial  
432 Communities and Metabolic Profiles of Chlamydia trachomatis Infection: Positioning Between  
433 Eubiosis and Dysbiosis. *Front Microbiol [Internet].* 2018 Mar 28 [cited 2018 Sep 24];9. Available  
434 from: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5883401/>
- 435 10. Vitali B, Cruciani F, Picone G, Parolin C, Donders G, Laghi L. Vaginal microbiome and metabolome  
436 highlight specific signatures of bacterial vaginosis. *Eur J Clin Microbiol Infect Dis.* 2015 Dec  
437 1;34(12):2367–76.
- 438 11. Gosmann C, Anahtar MN, Handley SA, Farcasanu M, Abu-Ali G, Bowman BA, et al. Lactobacillus-  
439 Deficient Cervicovaginal Bacterial Communities Are Associated with Increased HIV Acquisition in  
440 Young South African Women. *Immunity.* 2017 Jan 17;46(1):29–37.
- 441 12. Ratzke C, Gore J. Modifying and reacting to the environmental pH can drive bacterial interactions.  
442 *PLOS Biol.* 2018 Mar 14;16(3):e2004248.
- 443 13. Palmer RJ. Composition and development of oral bacterial communities. *Periodontol 2000*  
444 [Internet]. 2014 Feb [cited 2018 Sep 24];64(1). Available from:  
445 <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3876289/>
- 446 14. Tannock GW. A Special Fondness for Lactobacilli. *Appl Environ Microbiol.* 2004 Jun;70(6):3189–94.
- 447 15. Schmidt TSB, Hayward MR, Coelho LP, Li SS, Costea PI, Voigt AY, et al. Extensive transmission of  
448 microbes along the gastrointestinal tract. *Nieuwoudorp M, editor. eLife.* 2019 Feb 12;8:e42693.
- 449 16. Szajewska H, Ruszczyński M, Radzikowski A. Probiotics in the prevention of antibiotic-associated  
450 diarrhea in children: A meta-analysis of randomized controlled trials. *J Pediatr.* 2006 Sep  
451 1;149(3):367-372.e1.
- 452 17. Hempel S, Newberry SJ, Maher AR, Wang Z, Miles JNV, Shanman R, et al. Probiotics for the  
453 Prevention and Treatment of Antibiotic-Associated Diarrhea: A Systematic Review and Meta-  
454 analysis. *JAMA.* 2012 May 9;307(18):1959–69.
- 455 18. Ford AC, Quigley EMM, Lacy BE, Lembo AJ, Saito YA, Schiller LR, et al. Efficacy of Prebiotics,  
456 Probiotics, and Synbiotics in Irritable Bowel Syndrome and Chronic Idiopathic Constipation:  
457 Systematic Review and Meta-analysis. *Am J Gastroenterol.* 2014 Oct;109(10):1547–61.
- 458 19. Nikfar S, Rahimi R, Rahimi F, Derakhshani S, Abdollahi M. Efficacy of Probiotics in Irritable Bowel  
459 Syndrome: A Meta-Analysis of Randomized, Controlled Trials. *Dis Colon Rectum.* 2008 Dec  
460 1;51(12):1775–80.

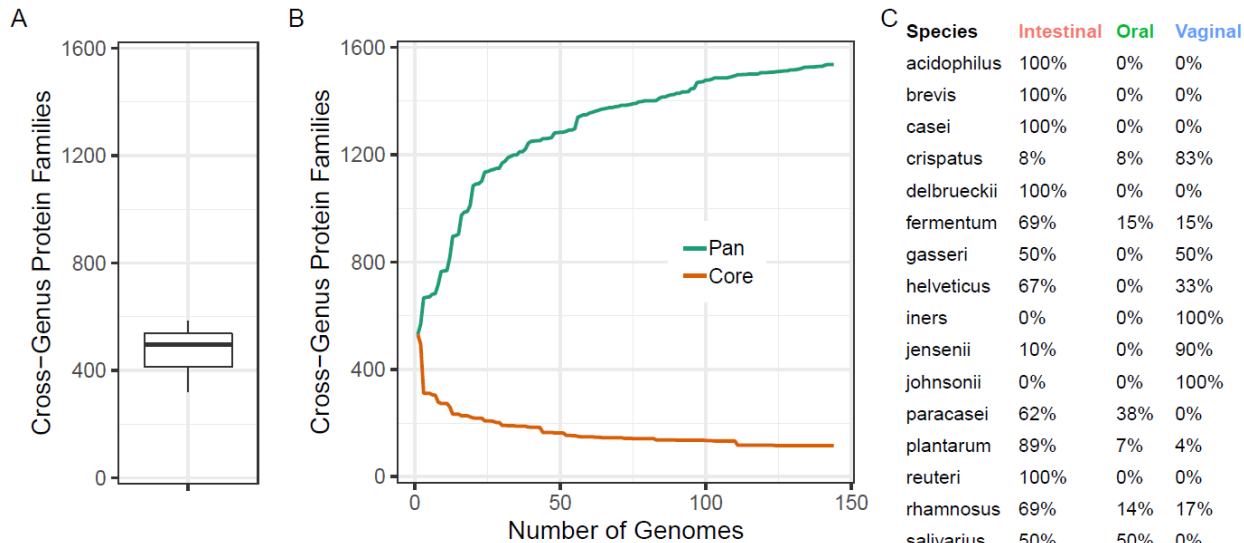
- 461 20. Elazab N, Mandy A, Gasana J, Vieira ER, Quizon A, Forno E. Probiotic Administration in Early Life,  
462 Atopy, and Asthma: A Meta-analysis of Clinical Trials. *Pediatrics*. 2013 Sep 1;132(3):e666–76.
- 463 21. Berstad A, Raa J, Midtvedt T, Valeur J. Probiotic lactic acid bacteria – the fledgling cuckoos of the  
464 gut? *Microb Ecol Health Dis [Internet]*. 2016 May 26 [cited 2018 Sep 24];27. Available from:  
465 <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4884264/>
- 466 22. Suez J, Zmora N, Zilberman-Schapira G, Mor U, Dori-Bachash M, Bashiardes S, et al. Post-Antibiotic  
467 Gut Mucosal Microbiome Reconstitution Is Impaired by Probiotics and Improved by Autologous  
468 FMT. *Cell*. 2018 Sep 6;174(6):1406-1423.e16.
- 469 23. de Vos WM. Lipotechoic acid in lactobacilli: D-alanine makes the difference. *Proc Natl Acad Sci*.  
470 2005;102(31):10763–4.
- 471 24. Granette C, Nutten S, Palumbo E, Morath S, Hermann C, Dewulf J, et al. Enhanced  
472 antiinflammatory capacity of a *Lactobacillus plantarum* mutant synthesizing modified teichoic acids.  
473 *Proc Natl Acad Sci*. 2005;102(29):10321–6.
- 474 25. Branco dos Santos F, de Vos WM, Teusink B. Towards metagenome-scale models for industrial  
475 applications—the case of Lactic Acid Bacteria. *Curr Opin Biotechnol*. 2013 Apr 1;24(2):200–6.
- 476 26. Le Barz M, Anhê FF, Varin TV, Desjardins Y, Levy E, Roy D, et al. Probiotics as Complementary  
477 Treatment for Metabolic Disorders. *Diabetes Metab J*. 2015 Aug;39(4):291–303.
- 478 27. Saulnier DM, Santos F, Roos S, Mistretta T-A, Spinler JK, Molenaar D, et al. Exploring Metabolic  
479 Pathway Reconstruction and Genome-Wide Expression Profiling in *Lactobacillus reuteri* to Define  
480 Functional Probiotic Features. *PLOS ONE*. 2011 Apr 29;6(4):e18783.
- 481 28. Lewis NE, Nagarajan H, Palsson BO. Constraining the metabolic genotype–phenotype relationship  
482 using a phylogeny of *in silico* methods. *Nat Rev Microbiol*. 2012 Apr;10(4):291–305.
- 483 29. Haggart CR, Bartell JA, Saucerman JJ, Papin JA. Whole-genome metabolic network reconstruction  
484 and constraint-based modeling. *Methods Enzymol*. 2011;500:411–33.
- 485 30. Kant R, Blom J, Palva A, Siezen RJ, de Vos WM. Comparative genomics of *Lactobacillus*. *Microb*  
486 *Biotechnol*. 2011 May;4(3):323–32.
- 487 31. Drissi F, Merhej V, Angelakis E, El Kaoutari A, Carrière F, Henrissat B, et al. Comparative genomics  
488 analysis of *Lactobacillus* species associated with weight gain or weight protection. *Nutr Diabetes*.  
489 2014 Feb;4(2):e109.
- 490 32. France MT, Mendes-Soares H, Forney LJ. Genomic Comparisons of *Lactobacillus crispatus* and  
491 *Lactobacillus iners* Reveal Potential Ecological Drivers of Community Composition in the Vagina.  
492 *Appl Env Microbiol*. 2016 Dec 15;82(24):7063–73.
- 493 33. Morita H, Toh H, Fukuda S, Horikawa H, Oshima K, Suzuki T, et al. Comparative Genome Analysis of  
494 *Lactobacillus reuteri* and *Lactobacillus fermentum* Reveal a Genomic Island for Reuterin and  
495 Cobalamin Production. *DNA Res*. 2008 Jun 1;15(3):151–61.

- 496 34. Zhang Z-G, Ye Z-Q, Yu L, Shi P. Phylogenomic reconstruction of lactic acid bacteria: an update. *BMC*  
497 *Evol Biol.* 2011 Jan 1;11:1.
- 498 35. Kleerebezem M, Vos WM de. Lactic acid bacteria: life after genomics. *Microb Biotechnol.* 2011 May  
499 1;4(3):318–22.
- 500 36. Rau MH, Zeidan AA. Constraint-based modeling in microbial food biotechnology. *Biochem Soc Trans.*  
501 2018 Mar 27;BST20170268.
- 502 37. Lewis NE, Hixson KK, Conrad TM, Lerman JA, Charusanti P, Polpitiya AD, et al. Omic data from  
503 evolved *E. coli* are consistent with computed optimal growth from genome-scale models. *Mol Syst*  
504 *Biol.* 2010 Jul 27;6:390.
- 505 38. Feist AM, Palsson BO. The biomass objective function. *Curr Opin Microbiol.* 2010 Jun;13(3):344–9.
- 506 39. Altafini C, Facchetti G. Metabolic Adaptation Processes That Converge to Optimal Biomass Flux  
507 Distributions. *PLoS Comput Biol.* 2015 Sep 4;11(9):e1004434.
- 508 40. Orth JD, Thiele I, Palsson BØ. What is flux balance analysis? *Nat Biotechnol.* 2010 Mar;28(3):245–8.
- 509 41. Pinto F, Medina DA, Pérez-Correa JR, Garrido D. Modeling metabolic interactions in a consortium of  
510 the infant gut microbiome. *Front Microbiol.* 2017;8:2507.
- 511 42. Schmidt BJ, Ebrahim A, Metz TO, Adkins JN, Palsson BØ, Hyduke DR. GIM3E: condition-specific  
512 models of cellular metabolism developed from metabolomics and expression data. *Bioinformatics.*  
513 2013 Nov 15;29(22):2900–8.
- 514 43. Li H, Zhu J. Targeted metabolic profiling rapidly differentiates *Escherichia coli* and *Staphylococcus*  
515 *aureus* at species and strain level. *Rapid Commun Mass Spectrom.* 2017;31(19):1669–76.
- 516 44. Benedict MN, Mundy MB, Henry CS, Chia N, Price ND. Likelihood-Based Gene Annotations for Gap  
517 Filling and Quality Assessment in Genome-Scale Metabolic Models. *PLOS Comput Biol.* 2014 Oct  
518 16;10(10):e1003882.
- 519 45. Thiele I, Vlassis N, Fleming RMT. fastGapFill: efficient gap filling in metabolic networks.  
520 *Bioinformatics.* 2014 Sep 1;30(17):2529–31.
- 521 46. Machado D, Andrejev S, Tramontano M, Patil KR. Fast automated reconstruction of genome-scale  
522 metabolic models for microbial species and communities. *Nucleic Acids Res.* 2018 Sep  
523 6;46(15):7542–53.
- 524 47. Devold S, Overbeek R, DeJongh M, Vonstein V, Best AaronA, Henry C. Automated Genome  
525 Annotation and Metabolic Model Reconstruction in the SEED and Model SEED. In: Alper HS, editor.  
526 *Systems Metabolic Engineering [Internet].* Humana Press; 2013 [cited 2017 Apr 6]. p. 17–45.  
527 (Methods in Molecular Biology). Available from: [http://dx.doi.org/10.1007/978-1-62703-299-5\\_2](http://dx.doi.org/10.1007/978-1-62703-299-5_2)
- 528 48. King B, Farrah T, Richards MA, Mundy M, Simeonidis E, Price ND. ProbAnnoWeb and ProbAnnoPy:  
529 probabilistic annotation and gap-filling of metabolic reconstructions. *Bioinformatics.* 2018 May  
530 1;34(9):1594–6.

- 531 49. Komatsuzaki N, Shima J, Kawamoto S, Momose H, Kimura T. Production of  $\gamma$ -aminobutyric acid  
532 (GABA) by *Lactobacillus paracasei* isolated from traditional fermented foods. *Food Microbiol.*  
533 2005;22(6):497–504.
- 534 50. Li H, Qiu T, Huang G, Cao Y. Production of gamma-aminobutyric acid by *Lactobacillus brevis* NCL912  
535 using fed-batch fermentation. *Microb Cell Factories.* 2010;9(1):85.
- 536 51. Walter J. Ecological Role of Lactobacilli in the Gastrointestinal Tract: Implications for Fundamental  
537 and Biomedical Research. *Appl Env Microbiol.* 2008 Aug 15;74(16):4985–96.
- 538 52. Valeur N, Engel P, Carbajal N, Connolly E, Ladefoged K. Colonization and immunomodulation by  
539 *Lactobacillus reuteri* ATCC 55730 in the human gastrointestinal tract. *Appl Environ Microbiol.*  
540 2004;70(2):1176–81.
- 541 53. Romero R, Hassan SS, Gajer P, Tarca AL, Fadrosh DW, Nikita L, et al. The composition and stability of  
542 the vaginal microbiota of normal pregnant women is different from that of non-pregnant women.  
543 *Microbiome.* 2014 Feb 3;2(1):4.
- 544 54. Gajer P, Brotman RM, Bai G, Sakamoto J, Schütte UME, Zhong X, et al. Temporal Dynamics of the  
545 Human Vaginal Microbiota. *Sci Transl Med.* 2012 May 2;4(132):132ra52-132ra52.
- 546 55. Ravel J, Gajer P, Abdo Z, Schneider GM, Koenig SSK, McCulle SL, et al. Vaginal microbiome of  
547 reproductive-age women. *Proc Natl Acad Sci.* 2011 Mar 15;108(Supplement 1):4680–7.
- 548 56. Al-Hassi HO, Mann ER, Sanchez B, English NR, Peake STC, Landy J, et al. Altered human gut dendritic  
549 cell properties in ulcerative colitis are reversed by *Lactobacillus plantarum* extracellular encrypted  
550 peptide STp. *Mol Nutr Food Res.* 2014;58(5):1132–43.
- 551 57. Bernardo D, Sánchez B, Al-Hassi HO, Mann ER, Urdaci MC, Knight SC, et al. Microbiota/Host  
552 Crosstalk Biomarkers: Regulatory Response of Human Intestinal Dendritic Cells Exposed to  
553 *Lactobacillus* Extracellular Encrypted Peptide. *PLOS ONE.* 2012 May 14;7(5):e36262.
- 554 58. Jewell JB, Kashket ER. Osmotically regulated transport of proline by *Lactobacillus acidophilus* IFO  
555 3532. *Appl Env Microbiol.* 1991 Oct 1;57(10):2829–33.
- 556 59. Vrancken G, Rimaux T, Weckx S, De Vuyst L, Leroy F. Environmental pH determines citrulline and  
557 ornithine release through the arginine deiminase pathway in *Lactobacillus fermentum* IMDO  
558 130101. *Int J Food Microbiol.* 2009 Nov 15;135(3):216–22.
- 559 60. Famularo G, Pieluigi M, Coccia R, Mastroiacovo P, Simone CD. Microecology, bacterial vaginosis and  
560 probiotics: perspectives for bacteriotherapy. *Med Hypotheses.* 2001 Apr 1;56(4):421–30.
- 561 61. Rousseau V, Lepargneur JP, Roques C, Remaud-Simeon M, Paul F. Prebiotic effects of  
562 oligosaccharides on selected vaginal lactobacilli and pathogenic microorganisms. *Anaerobe.* 2005  
563 Jun 1;11(3):145–53.
- 564 62. Witkin SS, Mendes-Soares H, Linhares IM, Jayaram A, Ledger WJ, Forney LJ. Influence of Vaginal  
565 Bacteria and d- and l-Lactic Acid Isomers on Vaginal Extracellular Matrix Metalloproteinase Inducer:

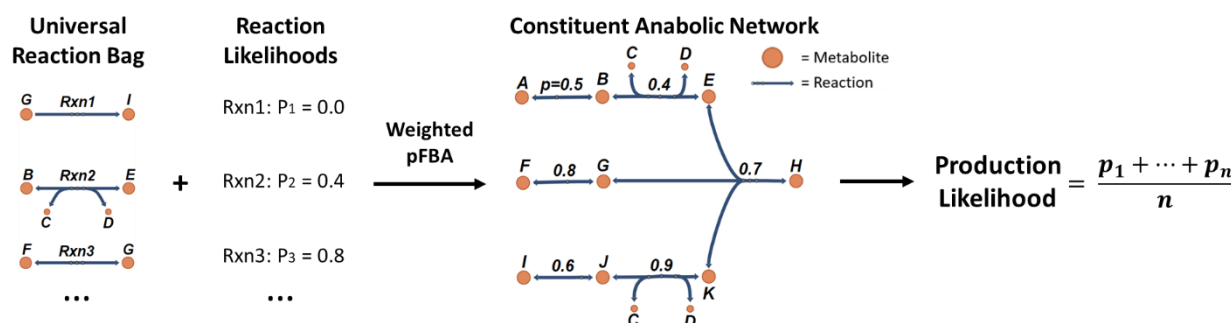
- 566        Implications for Protection against Upper Genital Tract Infections. *mBio*. 2013 Aug 30;4(4):e00460-  
567        13.
- 568        63. LeBlanc JG, Milani C, de Giori GS, Sesma F, van Sinderen D, Ventura M. Bacteria as vitamin suppliers  
569        to their host: a gut microbiota perspective. *Curr Opin Biotechnol*. 2013 Apr 1;24(2):160–8.
- 570        64. Henry CS, DeJongh M, Best AA, Frybarger PM, Linsay B, Stevens RL. High-throughput generation,  
571        optimization and analysis of genome-scale metabolic models. *Nat Biotechnol*. 2010 Sep;28(9):977–  
572        82.
- 573        65. Ebrahim A, Lerman JA, Palsson BO, Hyduke DR. COBRApy: COnstraints-Based Reconstruction and  
574        Analysis for Python. *BMC Syst Biol*. 2013 Aug 8;7(1):74.
- 575        66. Mundy M, Mendes-Soares H, Chia N. Mackinac: a bridge between ModelSEED and COBRApy to  
576        generate and analyze genome-scale metabolic models. *Bioinformatics*. 2017 Aug 1;33(15):2416–8.
- 577        67. Wegkamp A, Teusink B, De Vos W m., Smid E j. Development of a minimal growth medium for  
578        *Lactobacillus plantarum*. *Lett Appl Microbiol*. 2010 Jan 1;50(1):57–64.
- 579        68. Ricciardi A, Ianniello RG, Parente E, Zotta T. Modified chemically defined medium for enhanced  
580        respiratory growth of *Lactobacillus casei* and *Lactobacillus plantarum* groups. *J Appl Microbiol*. 2015  
581        Sep 1;119(3):776–85.
- 582        69. Elli M, Zink R, Rytz A, Reniero R, Morelli L. Iron requirement of *Lactobacillus* spp. in completely  
583        chemically defined growth media. *J Appl Microbiol*. 2000 Apr 1;88(4):695–703.
- 584        70. Rowland I, Gibson G, Heinken A, Scott K, Swann J, Thiele I, et al. Gut microbiota functions:  
585        metabolism of nutrients and other food components. *Eur J Nutr*. 2018 Feb 1;57(1):1–24.
- 586        71. Neis EPJG, Dejong CHC, Rensen SS. The Role of Microbial Amino Acid Metabolism in Host  
587        Metabolism. *Nutrients*. 2015 Apr 16;7(4):2930–46.
- 588        72. Wu G. Intestinal Mucosal Amino Acid Catabolism. *J Nutr*. 1998 Aug 1;128(8):1249–52.
- 589        73. Rooj AK, Kimura Y, Buddington RK. Metabolites produced by probiotic *Lactobacilli* rapidly increase  
590        glucose uptake by Caco-2 cells. *BMC Microbiol*. 2010 Jan 20;10(1):16.
- 591        74. Belzer C, Chia LW, Aalvink S, Chamlagain B, Piironen V, Knol J, et al. Microbial Metabolic Networks at  
592        the Mucus Layer Lead to Diet-Independent Butyrate and Vitamin B12 Production by Intestinal  
593        Symbionts. *mBio*. 2017 Nov 8;8(5):e00770-17.
- 594        75. Urrea V, Calle M. AUCRF: variable selection with random forest and the area under the curve. R  
595        Package Version 11. 2012;
- 596        76. Oksanen J, Blanchet FG, Kindt R, Legendre P, Minchin PR, O'hara R, et al. vegan: Community ecology  
597        package. R Package Version. 2011;117–8.
- 598        77. Ihaka R, Gentleman R. R: A Language for Data Analysis and Graphics. *J Comput Graph Stat*. 1996 Sep  
599        1;5(3):299–314.

- 600 78. Wickham H. *ggplot2: Elegant Graphics for Data Analysis*. Springer; 2016. 266 p.
- 601 79. Wickham H. *tidyr: Easily Tidy Data with spread () and gather () Functions*. Version 06 0. 2016;
- 602 80. Wickham H, Francois R, Henry L, Müller K. *dplyr: A grammar of data manipulation*. R Package  
603 Version 04. 2015;3.
- 604 81. Neuwirth E, Brewer RC. *ColorBrewer palettes*. R Package Version. 2014;1–1.
- 605 82. Haywood-Farmer E, Otto SP. The Evolution of Genomic Base Composition in Bacteria. *Evolution*.  
606 2003;57(8):1783–92.
- 607 83. Bentley SD, Parkhill J. Comparative Genomic Structure of Prokaryotes. *Annu Rev Genet*.  
608 2004;38(1):771–91.
- 609 84. Benjamini Y, Speed TP. Summarizing and correcting the GC content bias in high-throughput  
610 sequencing. *Nucleic Acids Res*. 2012 May;40(10):e72.
- 611 85. Gurevich A, Saveliev V, Vyahhi N, Tesler G. QUAST: quality assessment tool for genome assemblies.  
612 *Bioinformatics*. 2013 Apr 15;29(8):1072–5.
- 613
- 614



615

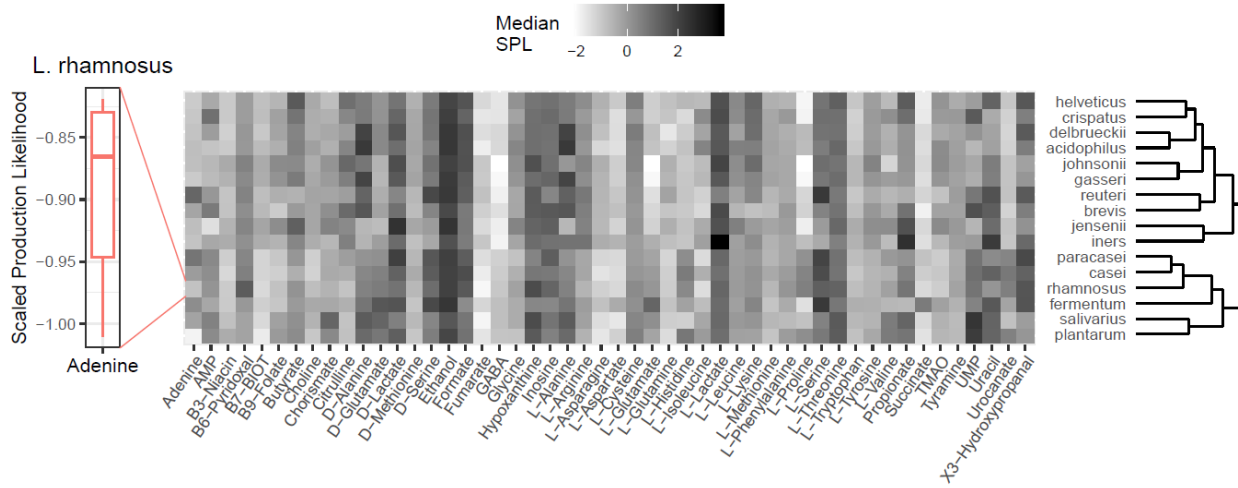
616 **Figure 1: Known metabolic annotations are extensively sampled across the 16 *Lactobacillus* species included in**  
 617 **this study.** The genomic features used for this analysis are PATRIC Cross-Genera Protein families (PGfams), a  
 618 standardized set of features across the PATRIC Database (4). (A) The number of metabolic PGfams for each  
 619 genome are shown here, with the median value indicated by the middle line in the boxplot. (B) For the 144 strains  
 620 from 16 species of *Lactobacillus*, we found that there are 116 protein families in the core set of metabolic PGfams,  
 621 while the pan set of PGfams expands to over 1500 families. The nearly plateau shape of the curve for the pan set  
 622 of PGfams curve indicates that this sampling represents a large portion of the genetic diversity among the 16  
 623 species included in the study. (C) This table shows the complete list of species used in this study and indicates the  
 624 percentage of strains that were isolated from each human body site. Each strain in this study is a member from  
 625 one of the 16 species and isolated from one of three human-associated body sites; intestinal, oral, or vaginal (Table  
 626 S2).



627

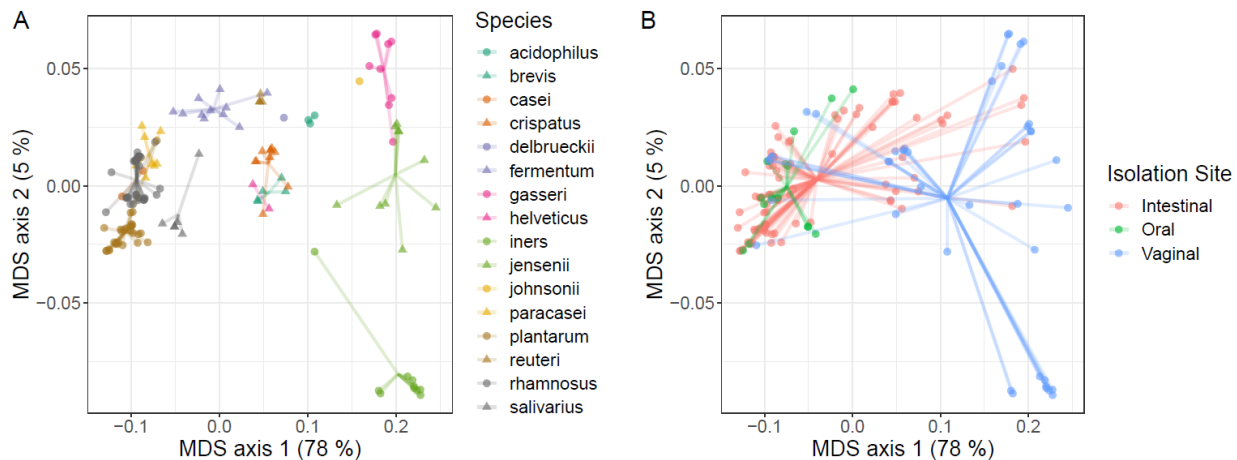
628 **Figure 2: PROTEAN is an approach for quantifying the likelihood that a given metabolic network, derived**  
 629 **exclusively from genomic evidence, is capable of synthesizing a particular metabolite.** A modified version of  
 630 Parsimonious Enzyme Usage FBA (weighted pFBA) was performed on a standardized set of reactions to generate  
 631 constituent anabolic networks for each genome. Reaction likelihoods were used to weight the minimization of flux  
 632 through each reaction in the network. Therefore, reactions with a greater likelihood were more likely to be  
 633 included in the resulting constituent anabolic network. Each constituent network has a set of input metabolites  
 634 representing the media condition (Table S4) and a demand reaction for a certain metabolic product. The resulting  
 635 constituent network is the set of reactions that requires flux to produce the metabolic product in the given media  
 636 condition. The production likelihood metric is an average of all the reaction likelihoods associated with the  
 637 reactions included in the constituent network. This metric is used as a summary statistic that allows for the

638 comparison of constituent networks across different metabolic products and strains, where a higher production  
 639 likelihood corresponds with greater genetic evidence for that particular constituent anabolic network.



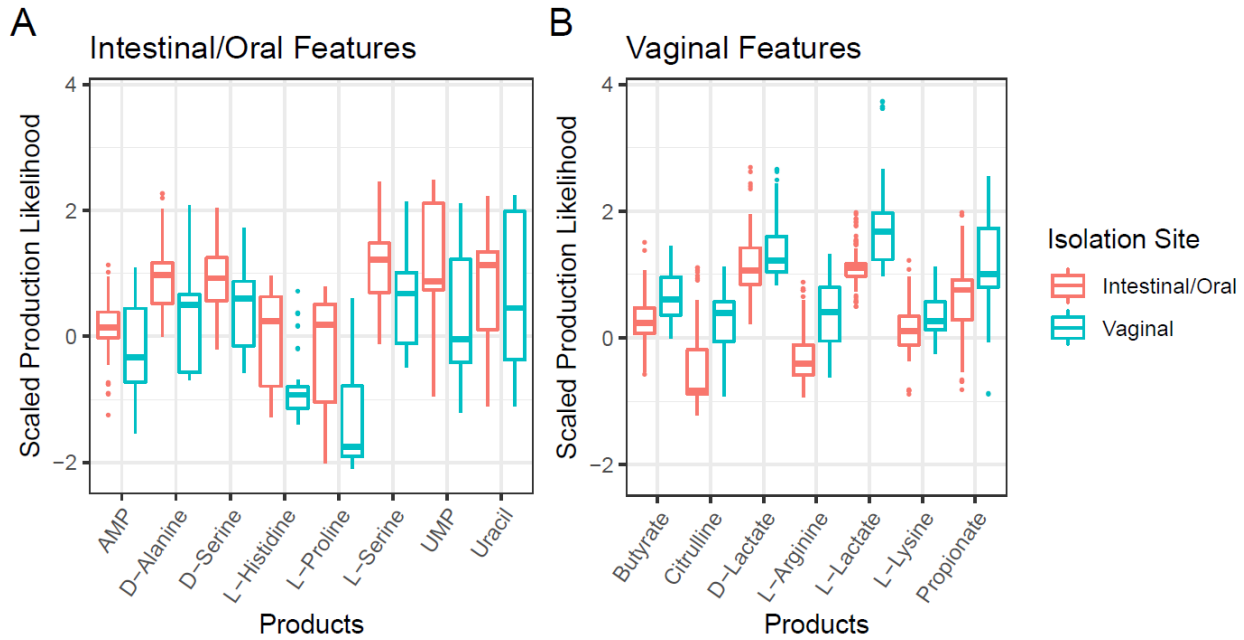
640

641 **Figure 3: Predicted metabolic production capabilities with the Scaled Production Likelihood (SPL) metric align**  
 642 **poorly with phylogeny.** There is a single production likelihood for each genome associated with each metabolite. A  
 643 median SPL can be calculated for a species that allows for more general comparisons across species, illustrated  
 644 here by the distribution for one species (*L. rhamnosus*) and one metabolite (adenine). There are 50 metabolites  
 645 used as features to allow for the comparison of predicted production capabilities across the lactobacilli analyzed.



646

647 **Figure 4: The Scaled Production Likelihood metric distinguishes metabolic functionality among species.** (A) We  
 648 found that *Lactobacillus* strains cluster significantly by species (PERMANOVA;  $P < 0.001$ ). (B) Additionally, they  
 649 cluster significantly by isolation site (PERMANOVA;  $P < 0.001$ ). Both plots are PCoA using the Bray-Curtis distance  
 650 metric of the SPLs for each isolate. Points in both panels are identical, but displayed with different color schemes.



652 **Figure 5: Machine learning of the SPL scores identifies metabolites that discriminate *Lactobacillus* strains.**  
653 Machine learning feature selection identified the metabolites that are both most likely to be produced by each  
654 group and capable of classifying the strains into two groups, intestinal/oral and vaginal, with greater than 90%  
655 accuracy. (A) There are eight metabolites that are more likely to be produced by the intestinal/oral isolates  
656 compared to the vaginal isolates. (B) There are seven metabolites that are more likely to be produced by the  
657 vaginal isolates compared to intestinal/oral isolates. Both models are more than 90% accurate in predicting the  
658 membership to which the given isolate belongs using the SPLs of the metabolites listed.

659