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MAGI: A method for metabolite, annotation, and gene integration

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15 **Author Contributions**

- OE, BPB, TRN conceived and designed the method 16
- 17 OE, BPB, OR, MT, TW, DWU developed the method
- KBL, MdR, CAH conducted the experiments 18
- OE, BPB, SD, TRN, KBL wrote the manuscript. 19
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23 Abstract

24

25 Metabolomics is a widely used technology for obtaining direct measures of 26 metabolic activities from diverse biological systems. However, ambiguous 27 metabolite identifications are a common challenge and biochemical 28 interpretation is often limited by incomplete and inaccurate genome-based 29 predictions of enzyme activities (*i.e.* gene annotations). Metabolite. 30 Annotation, and Gene Integration (MAGI) generates a metabolite-gene 31 association score using a biochemical reaction network. This is calculated by 32 a method that emphasizes consensus between metabolites and genes via 33 biochemical reactions. To demonstrate the potential of this method, we applied MAGI to integrate sequence data and metabolomics data collected 34 35 from Streptomyces coelicolor A3(2), an extensively characterized bacterium that produces diverse secondary metabolites. Our findings suggest that 36 37 coupling metabolomics and genomics data by scoring consensus between the 38 two increases the quality of both metabolite identifications and gene 39 annotations in this organism. MAGI also made biochemical predictions for 40 poorly annotated genes that were consistent with the extensive literature on 41 this important organism. This limited analysis suggests that using metabolomics data has the potential to improve annotations in sequenced 42 organisms and also provides testable hypotheses for specific biochemical 43 44 functions. MAGI is freely available for academic use both as an online tool at 45 https://magi.nersc.gov and with source code available at 46 https://github.com/biorack/magi.

47

48 Introduction

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50 Metabolomics approaches now enable global profiling, comparison, and 51 discovery of diverse metabolites present in complex biological samples¹. 52 Liquid chromatography coupled with electrospray ionization mass spectrometry (LC-MS) is one of the leading methods in metabolomics¹. A 53 critical measure in metabolomics datasets is known as a "feature," which is a 54 55 unique combination of mass-to-charge (m/z) and chromatographic retention time¹. Each distinct feature may match to hundreds of unique chemical 56 57 structures. This makes metabolite identification (the accurate assignment of the correct chemical structure to each feature) one of the fundamental 58 challenges in metabolomics²⁻⁴. To aid metabolite identification efforts, ions 59 (each with a unique m/z and retention time) are typically fragmented, and 60 the resulting fragments are compared against either experimental^{5, 6} or 61 computationally predicted^{5, 7-11} reference libraries. While this method is highly 62 effective at reducing the search space for metabolite identification, 63 64 misidentifications are inevitable, especially for metabolites lacking authentic 65 standards.

66

67 One strategy for addressing the large search space of compound 68 identifications is to assess identifications in the context of the predicted 69 metabolism of the organism(s) being studied. Several tools do this with 70 varying degrees of complexity with strategies including directly mapping 71 metabolites onto reactions¹² or scoring the likelihood of metabolite identities 72 using reaction networks and predictive pathway mapping¹³. However, many

metabolites cannot be included in these approaches. This is due to a number 73 of factors, including the low coverage in reaction databases^{14, 15} (especially for 74 secondary metabolites¹⁶⁻¹⁹), incomplete or inaccurate set of reactions for an 75 76 organism, and enzyme promiscuity not being taken into account when formulating the potential metabolism of an organism. To help address these 77 78 challenges computational strategies have been developed including 79 MyCompoundID ^{20, 21}, IIMDB ²², MINES ²³ and the ATLAS of biochemistry ²⁴ to 80 enzymatically enlarge compound space similar to the retrosynthesis tools 81 such as Retrorules ²⁵ and rePrime²⁶. These approaches can be complimented 82 by chemical networking to help address the limited number of metabolites 83 represented in reactions, by expanding reaction space based on chemical or 84 spectral similarity between metabolites. Effectively, even when a metabolite is not directly involved in a reaction, a linkage can still be made with a 85 reaction based on similarity to another well-studied metabolite^{16-19, 27}. In this 86 87 way, chemical networking is a viable solution that expands reaction 88 databases to integrate with already expansive metabolite databases. This 89 allows more putative metabolite identifications to be assessed using the 90 predicted metabolism of the organism(s).

91

92 Recently, approaches have been developed that span the gap between 93 metabolomics and genomics and allow for some enzyme promiscuity. GNP, 94 developed specifically for discovering new nonribosomal peptides (NRPs) and 95 polyketides, uses a gene-forward strategy that predicts possible chemical 96 structures of NRP and polyketide synthases and generates a set of predicted 97 MS/MS spectra based on those predictions; these predictions are then used to 98 mine MS data ²⁸. Pep2Path, also developed exclusively for NRPs and post-99 translationally modified peptides (RiPPs), takes a Bayesian approach to 100 scoring putative NRPs and RiPPs based on the gene sequences present in the assayed organism ²⁹. Finally, a more general approach has been developed 101 102 where a mutant library of an organism is assayed for major differences in the 103 mass spectrometry profile, and the major differences are manually annotated 104 with human intuition ³⁰.

105

106 Due to the vast amount of knowledge about Streptomyces species, they are 107 an excellent target for developing new tools for metabolite and genome 108 exploration. Representatives from this genus produce many antibacterials, 109 anticancer compounds, immunosuppresents, antifungals, cardiovascular 110 agents, and veterinary products including erythromycin, tetracvcline. doxorubicin, enediyenes, FK-506, rapamycin, avermectin, nemadectin, 111 112 amphotericin, griseofulvin, nystatin, lovastatin, compactin, monensin, and 113 tylosin ³¹. Thus making them a highly relevant group for in depth studies to link natural products with associated genes. In particular, Streptomyces 114 115 *coelicolor* is a model actinomycete secondary metabolite producer ³²; studies from over three thousand papers and over 60 years of work ³³ have produced, 116 117 among other things, a detailed understanding of the secondary metabolites this organism produces, where two are the pigmented antibiotics: 118 119 actinorhodin and undecylprodiosin. These experiments have identified the biochemical pathways, genes, and regulatory processes that are necessary 120 121 for producing the associated secondary metabolites ³⁴. 122

Here we report Metabolite, Annotation, and Gene Integration (MAGI), an 123 124 approach to generate metabolite-gene associations (Figure 1) by scoring consensus between metabolite identifications and gene annotations. MAGI is 125 126 guided by the principles that the probability of a metabolite identity increases 127 if there is genetic evidence to support that metabolite and that the 128 probability of a gene function increases if there is metabolomic evidence for 129 that function. Inputs to MAGI are typically a metabolite identification file of 130 LCMS features and a protein or gene sequence FASTA file. For each LCMS 131 feature, there are often many plausible metabolite identifications that can be 132 given a probability based on accurate mass error and/or mass fragmentation 133 comparisons. MAGI links these putative compound identifications to reactions both directly and indirectly by a biochemically relevant chemical similarity 134 135 network. Likewise, MAGI associates input sequences to biochemical reactions 136 by assessing sequence homology to reference sequences in the MAGI 137 reaction database. For each sequence, there are often several plausible 138 reactions with equal or similar probability. While annotation services would 139 typically reduce specificity in these cases (e.g., by simply annotating as 140 oxidoreductase), MAGI maintains all specific reactions as possibilities. Since 141 MAGI links both metabolites and sequences to reactions with numerical 142 scores that are proxies for probabilities, a final integrative MAGI score is 143 calculated that magnifies consensus between a gene annotation and a 144 metabolite identification. We applied this approach to one of the best 145 characterized secondary metabolite producing bacteria, Streptomyces 146 *coelicolor* A3(2)³⁵, by integrating its genome sequence with untargeted 147 metabolomics data. MAGI successfully reduced the metabolite identity search 148 space by scoring metabolite identities based on the predicted metabolism of 149 an organism. Additionally, further investigation of the metabolite-gene 150 associations led to identification of unannotated and misannotated genes that were subsequently validated using literature searches. This simple 151 152 example illustrates the key aspects of MAGI.

153

154 Methods

155

156 Media and culture conditions. A 20 µL volume of glycerol stock of wild-157 type *S. coelicolor* spores was cultured in 40 mL R5 medium in a 250-mL flask. 158 One liter of R5 medium base included 103 g sucrose, 0.25 g K_2SO_4 , 10.12 g 159 MgCl₂•6H₂0, 10 g glucose, 0.1 g cas-amino acids, 2 mL trace element 160 solution, 5 g yeast extract, and 5.73 g TES buffer to 1 L distilled water. After autoclave sterilization, 1 mL 0.5% KH₂PO₄, 0.4 mL 5M CaCl₂•2H₂0, 1.5 mL 161 20% L-proline, 0.7 ml 1N NaOH were added as per the following protocol: 162 163 https://www.elabprotocols.com/protocols/#!protocol=486. Each flask 164 contained a stainless steel spring (McMaster-Carr Supply, part 9663K77), cut 165 to fit in a circle in the bottom of the flask. The spring was used to prevent 166 clumping of S. coelicolor during incubation. A foam stopper was used to close 167 each flask (Jaece Industries Inc., Fisher part 14-127-40D). Four replicates of 168 each sample were grown in a 28°C incubator with shaking at 150 rpm. On 169 day six, 1 mL from each replicate were collected in 2 mL Eppendorf tubes in a 170 sterile hood. Samples were centrifuged at 3,200 x g for 8 minutes at 4 °C to 171 pellet the cells. Supernatants were decanted into fresh 2 mL tubes and frozen 172 at -80 °C. Pellets were flash frozen on dry ice and then stored at -80 °C.

173

174 **LCMS sample preparation and data acquisition.** In preparation for LCMS, medium samples were lyophilized. Dried medium was then extracted with 175 176 150 µL methanol containing an internal standard (2-Amino-3-bromo-5-177 methylbenzoic acid, 1 µg/mL, Sigma, #631531), vortexed, sonicated in a 178 water bath for 10 minutes, centrifuged at 5,000 rpm for 5 min, and 179 supernatant finally centrifuge-filtered through a 0.22 µm PVDF membrane 180 (UFC40GV0S, Millipore). LC-MS/MS was performed in negative ion mode on a 181 2 µL injection, with UHPLC reverse phase chromatography performed using 182 an Agilent 1290 LC stack and Agilent C18 column (ZORBAX Eclipse Plus C18, 183 Rapid Resolution HD, 2.1 x 50 mm, 1.8 μ m) at 60 °C and with MS and MS/MS 184 data collected using a QExactive Orbitrap mass spectrometer (Thermo 185 Scientific, San Jose, CA). Chromatography used a flow rate of 0.4 mL/min, first equilibrating the column with 100% buffer A (LC-MS water with 0.1% formic 186 187 acid) for 1.5 min, then diluting over 7 minutes to 0% buffer A with buffer B 188 (100% acetonitrile with 0.1% formic acid). Full MS spectra were collected at 189 70,000 resolution from m/z 80-1,200, and MS/MS fragmentation data 190 collected at 17,500 resolution using an average of 10, 20 and 30 eV collision 191 energies.

192

193 Feature detection. MZmine (version 2.23) ³⁶ was used to deconvolute mass 194 spectrometry features. The methods and parameters used were as follows (in 195 the order that the methods were applied). MS/MS peaklist builder: retention 196 time between 0.5-13.0 minutes, m/z window of 0.01, time window of 1.00. 197 Peak extender: m/z tolerance 0.01 m/z or 50.0 ppm, min height of 1.0E0. 198 Chromatogram deconvolution: local minimum search algorithm where 199 chromatographic threshold was 1.0%, search minimum in RT range was 0.05 200 minutes, minimum relative height of 1.0%, minimum absolute height of 1.0E5, minimum ratio of peak top/edge of 1.2, peak duration between 0.01 201 202 and 30 minutes. Duplicate peak filter: m/z tolerance of 0.01 m/z or 50.0 ppm, 203 RT tolerance of 0.15 minutes. Isotopic peaks grouper: m/z tolerance of 1.0E-6 204 m/z or 20.0 ppm, retention time tolerance of 0.01, maximum charge of 2, 205 representative isotope was lowest m/z. Adduct search: RT tolerance of 0.01 206 minutes, searching for adducts M+Hac-H, M+Cl, with an m/z tolerance of 207 1.0E-5 m/z or 20.0 ppm and max relative adduct peak height of 1.0%. Join aligner: m/z tolerance of 1.0E-6 m/z or 50.0 ppm, weight for m/z of 5, 208 209 retention time tolerance of 0.15 minutes, weight for RT of 3. Same RT and m/ 210 z range gap filler: *m/z* tolerance of 1.0E-6 *m/z* or 20.0 ppm.

211

212 Metabolite identification. During the LCMS acquisition, two MS/MS 213 spectra were acquired for every MS spectrum. These MS/MS spectra are 214 acquired using data-dependent criteria in which the 2 most intense ions are 215 pursued for fragmentation, and then the next 2 most intense ions such that 216 no ion is fragmented more frequently than every 10 seconds. To assign 217 probable metabolite identities to a spectrum a modified version of the 218 previously described MIDAS approach was used⁷. Our metabolite database is 219 the merger of HMDB, MetaCyc, ChEBI, WikiData, GNPS, and LipidMaps resulting in approximately 180,000 unique chemical structures. For each of 220 221 these structures, a comprehensive fragmentation tree was pre-calculated to 222 a depth of 5 bond-breakages; these trees were used to accelerate the MIDAS

scoring process. The source code to generate trees and score spectra 223 224 against trees is available on GitHub (https://github.com/biorack/pactolus). 225 The following procedure was used in the MIDAS scoring. Precursor m/z values 226 were neutralized by 1.007276 Da. For each metabolite within 10 ppm of the 227 neutralized precursor mass, MS/MS ions were associated with nodes of the fragmentation tree using a window of 0.01 Da using MS/MS neutralizations of 228 229 1.00727, 2.01510, and -0.00055, as described ⁷. For metabolite-features of 230 interest discussed in the text, retention time, m/z, adduct, and fragmentation 231 pattern were used to define a Metabolite Atlas ³⁷ library (Supplementary Data 232 1). For each metabolite, raw data was inspected manually using MZmine 36 to 233 rule out peak misidentifications due to adduct formation and in-source 234 degradation.

235 MAGI biochemical reaction and reference sequence database. The 236 MAGI biochemical reaction database was constructed by aggregating all 237 publicly available biochemical reactions in MetaCyc and RHEA biochemical 238 reaction databases ^{14, 15}. This reaction database currently includes 12,293 239 unique metabolite structures. Identical reactions were collapsed together by 240 calculating a "reaction InChI key," where the SMILES strings of all members 241 of a reaction were joined together, separated by a "." and converted to a 242 single InChI string through an RDkit (https://github.com/rdkit/rdkit) Mol 243 object, and then the InChI key was calculated also using RDKit. Biochemical reactions with identical reaction InChI keys have identical chemical 244 245 metabolites, indicating they are duplicates, and were collapsed into one 246 database entry, retaining reference sequences. Reference sequences for 247 each biochemical reaction from each database were combined to create a set 248 of curated reference sequences for each biochemical reaction in the 249 database.

250

251 **Chemical Network.** In order to expand the chemical space beyond what is 252 in the biochemical reaction database, a chemical network was constructed to 253 relate all metabolites in the database to metabolites in biochemical reactions 254 by biochemical similarity. In each molecule, 70 chemical features These features were defined 255 (Supplementary Table 1) were located. previously as being biochemically relevant ³⁸. The count of each feature was 256 257 stored as a vector for each molecule. The Euclidean distance between two 258 vectors was used to determine similarity between two molecules and 259 construct a similarity network where every molecule is connected to every 260 molecule by the difference in their vectors. This network was trimmed by 261 calculating a minimum-spanning tree based on frequency of biochemical 262 differences where more frequent differences would be preserved when 263 possible (Supplementary Data 2).

264

Gene Annotations of Streptomyces coelicolor. KEGG annotations were obtained by submitting the S. coelicolor protein FASTA obtained from IMG to the KEGG Automatic Annotation Server version 2.1 ³⁹ and downloading the gene-KO results table. KO numbers were associated with reactions by assessing if there was a link to one or more KEGG reaction entries directly from the webpage of that KO. For BioCyc annotations and reactions, the BioCyc *S. coelicolor* database was downloaded. For the reactions in Table 1, KEGG and BioCyc reactions were manually inspected and compared to MAGIreactions.

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275 MAGI workflow. An input metabolite structure is expanded to similar 276 metabolite structures as suggested by the chemical network and all 277 tautomers of those metabolites. Searching all tautomeric forms of a 278 metabolite structure is a known method to enhance metabolite database 279 searches ⁴⁰. Tautomers were generated by using the MolVS package. The 280 reaction database is then queried to find reactions containing these 281 metabolites or their tautomers. Direct matches are stereospecific, but 282 tautomer matches are not. This is due to limitations in the tautomer 283 generating method and in how the chemical network was constructed. The 284 metabolite score, C, is inherited from the MS/MS scoring algorithm and is a 285 proxy for the probability that a metabolite structure is correctly assigned. In 286 our case, it is the MIDAS score, but could be any score due to the use of the 287 geometric mean to calculate the MAGI score. The metabolite score is set to 1 288 as a default.

289

290 If the reaction has a reference sequence associated with it, this reference 291 sequence is used as a BLAST guery against a sequence database of the input 292 gene sequences to find genes that may encode that reaction. The reciprocal 293 BLAST is also performed, where genes in the input gene sequences are 294 queries against the reaction reference sequence database; this finds the 295 reactions that a gene may encode for. The BLAST results are joined by their 296 common gene sequence and are used to calculate a homology score: 297 H=F+R-|F-R| where F and R are log-transformed e-values of the BLAST 298 results (a proxy for the probability that two gene sequences are homologs), 299 with F representing the reaction-to-gene BLAST score, and R the gene-to-300 reaction BLAST score. The homology score is set to 1 if no sequence is 301 matched.

302

303 The reciprocal agreement between both BLAST searches is also assessed, 304 namely whether they both agreed on the same reaction or not, formulating a 305 reciprocal agreement score: α . α is equal to 2 for reciprocal agreements, 1 for 306 disagreements that had BLAST score within 75% of the larger score, 0.01 for 307 disagreements with very different BLAST scores, and 0.1 for situations where 308 one of the BLAST searches did not yield any results. For cases where 309 metabolites are linked to reactions but there is not a reference protein 310 sequence available, a weight factor, X, is needed. We chose, X, such that: i) X=0.01 when a metabolite is not in any reaction; ii) X=1.01 when a 311 312 metabolite is in reaction missing a reference sequence; and iii) X=2.01 when 313 a metabolite is in a reaction with a sequence. These arbitrary scores were selected solely to distinguish between different "agreement" states during 314 315 the reciprocal BLAST. We did not observe much difference in the plurality of 316 compound annotations depending on these weights (data not shown), however, they did have an impact on the number of annotations that agreed 317 318 with KEGG and MetaCyc (Supplementary Figure 4). The most impactful 319 weight appeared to be the "close reciprocal disagreement," meaning that 320 there was not an exact match in the bidirectional BLAST, but the e-scores

were within the given threshold. If this weight was low (0.01) or high (2.0),
 there were fewer annotations that agreed with KEGG and MetaCyc.

323

The final MAGI-score $M = GM[[C, H, a, X]]/n^{L}$ is a proxy for the probability that 324 a gene and metabolite are associated. M is generated by calculating the 325 326 geometric mean (GM) of the metabolite score (C), homology score (H), 327 reciprocal agreement score (α) and weight factor (X), and whether or not the 328 metabolite is present in a reaction (n^{L}) where L is the network level 329 connecting the metabolite to a reaction (a proxy for the probability that a 330 compound is involved in a reaction) and *n* is a penalty factor for the network 331 level. Currently, n is equal to 4, but this parameter may change as the 332 scoring function is optimized and more training data is acquired. We did not 333 observe this penalty factor to greatly affect the number of gene annotations 334 that agreed with KEGG or MetaCyc, though this did have a large impact on 335 the number of features with multiple suggestions for compound identities; 336 the higher the penalty factor, the lower the number of compound 337 suggestions. MAGI often gives a high score to multiple metabolites, which is 338 not surprising given the relatedness of many metabolites (e.g. isomers). 339 Therefore, we recommend carefully considering the top scoring molecules 340 and not assuming that the top ranked one is correct (Supplementary Figure 341 5). Additional benchmarking analysis shows agreement between KEGG, 342 BioCyc and MAGI annotations for high MAGI homology scores (Supplementary 343 Discussion and SI Figures 2 and 3). The geometric mean was used to account 344 for the different scales of the individual scores, but weights may be applied to 345 each individual score during the geometric mean calculation to further fine-346 tune the MAGI scoring process. We expect the weights to become further 347 optimized as more results are processed through MAGI.

348

349 The final output is a table representing all unique metabolite-reaction-gene 350 associations, their individual scores, and their integrated MAGI score 351 (Supplementary Table 2). For scoring metabolite identities, a slice of this final 352 output is created by retaining the top scoring metabolite-reaction-gene 353 association for each unique metabolite structure; these can be mapped back 354 onto the mass spectrometry results table to aid the identification of each 355 mass spectrometry feature. For assessing gene functions, another slice of this final output is created by retaining the top scoring metabolite-reaction-356 357 gene association for each unique gene-reaction pair. For a typical bacterial 358 genome of \sim 6000 genes and a metabolites file of \sim 6000 compounds, the 359 MAGI calculation performed via the web service at https://magi.nersc.gov/ 360 should take about thirty minutes to complete. While MAGI can provide 361 valuable insights into primary metabolism, these reactions tend to be better 362 characterized and therefore a particularly important application of MAGI is for 363 secondary metabolite pathways.

364

365 Data Availability

All source code is available at <u>https://github.com/biorack/magi</u>, and the *S. coelicolor* mass spectrometry data (.mzML files) and MIDAS results
(metabolite_0ae82b08.csv) can be found here: <u>https://magi.nersc.gov/jobs/?</u>
<u>id=0ae82b08-b2a3-40d8-bb9a-e64b567cacd2</u>.

371 Application Availability and Usage

372 Potential MAGI users may use the application on their personal computers by 373 downloading the source code from the GitHub repository, or may upload their 374 data files to the web service. In order to use MAGI, users must provide at 375 least one of the following: a FASTA file of genes they wish to be associated to reactions, and/or a metabolites file they wish to associate to reactions. The 376 377 metabolites file should be in a table file format (e,q, .csv, .tsv, Excel), and 378 must have a column named "original compound" that describes the InChI 379 Key for each metabolite of interest. If both FASTA and metabolite files are 380 provided, then associations between genes and metabolites will be made as 381 well.

382

383 Results and Discussion384

385 Improved metabolite identification for metabolomics. To examine how 386 MAGI uses genomic information to filter and score possible metabolite 387 identities from a metabolomics experiment, sequencing and metabolomics 388 data were obtained for S. coelicolor. After processing the raw LCMS data to 389 find chromatograms and peaks, 878 features with a unique m/z and retention 390 time were found in the dataset. After neutralizing the m/z values, accurate 391 mass searching, and conducting MS/MS fragmentation pattern analysis, 6,604 392 unique metabolite structures were tentatively associated with these features 393 (Supplementary Table 3). This means on average there were almost 8 394 candidate structures for each feature. For a candidate structure to be 395 associated with a feature, it must have at least one matching fragmentation 396 spectrum. As this is often the method for identifying metabolites, it highlights 397 the problem in deconvolution of a signal to a specific chemical structure. 398 2,786 of these structures were then linked to a total of 10,265 reactions 399 either directly or via the chemical similarity network, and the reactions were 400 associated with 3,181 (out of 8,210) S. coelicolor genes by homology. Finally, 401 a MAGI score was calculated for each metabolite-reaction-gene association 402 (Supplementary Table 4).

403

404 An example that illustrates MAGI's utility in determining the most likely 405 correct metabolite identification is the feature putatively identified as 1,4-406 dihydroxy-6-naphthoic acid. Here, a feature with an m/z of 203.0345 was 407 observed. This feature was associated with the chemical formula $C_{11}H_8O_4$, 408 which could be derived from 16 unique chemical structures in the metabolite 409 database (Supplementary Table 5). Mass fragmentation spectra were 410 collected for this feature and analyzed using MIDAS⁷, a tool that scores the 411 observed fragmentation spectrum against its database of in-silico 412 fragmentation trees for the 16 potential structures. Based only on the MIDAS 413 5,6-dihydroxy-2metabolite score, the top scoring structure was methylnaphthalene-1,4-dione. However, after calculating the MAGI scores, a 414 415 different metabolite received the highest score. Of the 16 potential 416 metabolites, only 1,4-dihydroxy-6-naphthoic acid was in a reaction that had a 417 perfect match to genes in S. coelicolor (an E-value of 0.0 to SCO4326; Table 1). This metabolite is a known intermediate in an alternative menaguinone 418 biosynthesis pathway discovered in *S. coelicolor*^{41, 42}, making it much more 419

420 likely to be a metabolite detected from the metabolome of *S. coelicolor* as421 opposed to the metabolite found by looking at mass fragmentation alone.

422

423 Metabolomics-driven gene annotations. MAGI keeps the biochemical 424 potential of an organism unconstrained by considering a plurality of probable 425 gene product functions. One effect of this is that more reactions are 426 associated with genes than other services (Figure 2A). Because reactions are 427 the pivotal link between metabolites and genes, this allows integration of a 428 larger fraction of a metabolomics dataset with genes. Furthermore, MAGI 429 associates many genes that are not annotated using traditional approaches 430 with at least one reaction (Figure 2B). Out of a total of 8,210 predicted coding sequences in S. coelicolor, KEGG and BioCyc have one or more reactions 431 432 associated with 1,106 and 1,294 genes, respectively. On the other hand, 433 MAGI associated 5,209 genes with one or more reactions, out of which 3,719 434 genes had no reaction associated with them in either KEGG or BioCyc (Figure 435 2B). Of these 3,719 genes, 1,883 were linked to at least one metabolite in the 436 metabolomics data (Supplementary Table 4). Certainly, not all MAGI gene-437 reaction associations are correct, however, this does provide many testable 438 hypotheses that give footholds to discover new biochemistry As can be seen 439 in Figure 2C, many of these new gene-reaction associations have high scores, 440 indicating a likely connection.

441

442 Validation of gene-metabolite integration in pathways. One of the 443 most well-known biosynthetic pathways in *S. coelicolor* is the pathway to synthesize the pigmented antibiotic actinorhodin³⁵. We examined the MAGI 444 445 results involving the metabolites and genes of actinorhodin biosynthesis as a 446 proof-of-principle that MAGI successfully integrates metabolites and genes, 447 and that these results can be mapped onto a reaction network. Actinorhodin and all of its detected intermediates were correctly identified and accurately 448 449 mapped to the correct genes (Figure 3A), despite some intermediates having 450 several plausible metabolite identities (Supplementary Table 6). Notably, KEGG did not annotate the majority of actinorhodin biosynthesis genes, and 451 452 the one gene that it did annotate was incorrect (Table 1).

453

454 In another example, we examined the menaguinone biosynthesis pathway, which is essential for respiration in bacteria⁴³ and thus should be included in 455 456 every metabolic reconstruction for organisms that produce menaquinone. An 457 alternative menaguinone biosynthesis pathway was recently discovered and validated in *S. coelicolor*^{41, 42}, serving as another proof-of-principle exercise for 458 459 assessing the MAGI platform. MAGI linked 4 of 7 intermediate metabolites of 460 the pathway to the appropriate genes (Figure 3B, Supplementary Table 7). 461 Interestingly, while KEGG accurately assigned reactions to all but one of the 462 genes in this biosynthetic pathway, BioCyC had vague textual annotations and no reactions (Table 1). Therefore, a metabolomics tool that relies on 463 464 BioCyc model for S. coelicolor would be unable to integrate any of these 465 metabolites with genes for the purpose of either improved metabolite 466 identifications or gene annotations.

467

468 **Correction of annotation errors.** Gene annotation pipelines are 469 notoriously error-prone⁴⁴ and yield inconsistent results based on the

bioinformatic analyses used: the database used for homology searches, and 470 what kind of additional data (e.g. PFams, genetic neighborhoods, and 471 472 literature mining) are incorporated into the annotation algorithm or not (see 473 Table 1 for some examples). For example, the undecylprodigiosin synthase gene is known⁴⁵, yet was incorrectly annotated in the KEGG genome 474 annotation for S. coelicolor. KEGG annotated this gene as "PEP utilizing 475 476 enzyme" with an EC number of 2.7.9.2 (pyruvate, water phosphotransferase 477 with paired electron acceptors). This error is notable because the 478 undecylprodigiosin synthase reaction has an EC number of 6.4.1.-: ligases 479 that form carbon-carbon bonds. On the other hand, BioCyc correctly 480 annotates SCO5896 as undecylprodigiosin synthase, presumably using 481 manual curation or a thorough literature-searching algorithm.

482

483 MAGI used metabolomics data to score the possible gene annotations for 484 SCO5896 in addition to homology scoring (*i.e.* E-value). In the absence of 485 metabolomics data, MAGI initially associated the SCO5896 gene sequence 486 with the prodigiosin synthase and norprodigiosin synthase reactions via 487 BLAST searches against the MAGI reaction reference sequence database 488 (Figure 4). Metabolomics analysis revealed that the feature with an m/z of 489 392.2720 could potentially be undecylprodigiosin, which MAGI associated 490 with only the undecylprodigiosin synthase reaction (Figure 4). Because this 491 reaction does not have a reference sequence in our database, it could not be 492 queried against the S. coelicolor genome. However, the chemical network 493 revealed that prodigiosin is a similar metabolite that is in a reaction that does 494 have a reference sequence (Figure 4). When the prodigiosin synthase 495 reaction's reference sequence was gueried against the *S. coelicolor* genome, 496 the top hit was SCO5896, thus making a reciprocal connection between the 497 mass spectrometry feature and gene via the prodigiosin synthase reaction 498 (Figure 4).

499

500 Making nonexistent or vague annotations specific. The vast majority of 501 sequenced genes have no discrete functional predictions, preventing the in-502 depth understanding of metabolic processes of most organisms. S. coelicolor is well known to produce several polyketides and is known to have the 503 504 genetic potential to produce many more. The SC05315 gene product is WhiE, a known polyketide aromatase involved in the biosynthesis of a white 505 pigment characteristic of S. coelicolor^{46, 47}. KEGG and BioCyC textually 506 507 annotated the gene as "aromatase" or "polyketide aromatase," but neither links the gene to a discrete reaction. Although the text annotations are 508 509 correct, the lack of a biochemical reaction prohibits the association of this 510 gene with metabolites. On the other hand, MAGI was successfully able to 511 associate SCO5315 with an observed metabolite (20-carbon polyketide 512 intermediate with an m/z of 401.0887) via a polyketide cyclization reaction with a MAGI consensus score of 4.59 (Table 1). While the physiological 513 514 function of WhiE is to cyclize a 24-carbon polyketide intermediate, the 515 enzyme has been shown to also catalyze the cyclization of similar polyketides with varying chain length, including the 20-carbon species observed in the 516 metabolomics data presented here⁴⁸⁻⁵⁰. 517

In another example where other annotation services were unable to assign 519 520 any reactions to a gene product, MAGI associated SCO7595 with the anhydro-NAM kinase reaction via the detected metabolite anhydro-N-acetylmuramic 521 522 acid (anhydro-NAM) (m/z 274.0941) (Table 1). Anhydro-NAM is an intermediate in bacterial cell wall recycling, a critically important and 523 524 significant metabolic process in actively growing bacterial cells; E. coli and 525 other bacteria were observed to recycle roughly half of cell wall components per generation^{51, 52}. MAGI also associated anhydro-NAM to SCO6300 via an 526 527 acetylhexosaminidase reaction (Table 1) that produces the metabolite. KEGG 528 and RAST both annotate this gene to be acetylhexosaminidase with a total of 529 5 possible reactions, but none involve anhydro-NAM (Table 1). The detection 530 of anhydro-NAM may be considered orthogonal experimental evidence to 531 indicate that SCO6300 can act on N-acetyl-B-D-glucosamine-anhydro-NAM along with the other acetylhexosamines predicted by KEGG and RAST, 532 forming an early stage in anhydromurpoeptide recycling. In the absence of 533 534 MAGI, a researcher may have been able to manually curate a metabolic 535 model by manually assessing the text annotations and adding reactions to 536 the model, but the MAGI framework not only makes this process easier, it 537 also connects an experimental observation that supports the predicted 538 function of the gene.

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540 Potential for making novel annotations. In addition to these few 541 examples, there are hundreds more gene-reaction-metabolite associations 542 that could be used to strengthen, validate, or correct existing annotations from KEGG or BioCyc, as well as discover new annotations through 543 experimentation. These MAGI associations can be sorted by their MAGI score 544 to generate a ranked list of candidate genes and gene functions, with 545 546 optional hierarchical grouping and filtering of the list by homology, metabolite, chemical network, and/or reciprocal score. For example, of the 547 548 1,883 S. coelicolor genes that were uniquely linked to a metabolite via a 549 reaction by MAGI, roughly one-third were connected directly to a metabolite; 550 that is, the chemical similarity network was not used to expand reaction 551 space (Figure 5A and Figure 2C teal markers). Furthermore, one-third of 552 these genes had perfect reciprocal agreement between the metabolite-to-553 gene and gene-to-metabolite search directions (Figure 5B and Figure 2C teal 554 circles). These 190 genes can be further separated or binned based on their 555 homology score or MAGI score (Figure 5C), resulting in an actionable number 556 of high-priority and high-strength novel gene function hypotheses to test in 557 future studies.

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559 Limitations of this study. In this study, we show that MAGI produces 560 plausible associations between genes and metabolites from Streptomyces coelicolor. Since the associations shown in this paper are judged by manual 561 562 inspection, there are not enough validated links to compute a reliable false 563 discovery rate or applicability to other systems. Therefore an important 564 future work will be to broadly apply MAGI across many organisms and 565 evaluate the generality of this approach. This will ensure that the parameters 566 used are not over fit specifically to Streptomyces coelicolor. In addition, given the paucity of direct biochemical validations of gene functions, it will likely be 567 568 necessary to integrate MAGI with high throughput mutagenesis studies to

accurately determine false discovery rates. Lastly, more unique metabolites
can be observed by combining data collected from polar and lipid fractions of
metabolites along with combining positive and negative ionization modes.
The results here are based on measured signals from a small subset of the
Streptomyces coelicolor metabolome.

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576 **Conclusion**

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578 In this work we describe MAGI, a method for integrating metabolomics 579 observations with genomic predictions to help overcome the limitations of 580 each and strengthen the biological conclusions made by both. Using 581 Streptomyces coelicolor as a test case, we find that this method can help 582 strengthen metabolite identifications, suggests specific biochemical 583 predictions about genes that may otherwise be ambiguous, and suggests new biochemistry via the chemical network. It will be important to also 584 585 evaluate this approach for diverse organisms to determine the generality of the method. In order to facilitate broad usage by the academic community, 586 587 we provide MAGI through the National Energy Research Scientific Computing 588 Center (NERSC) at https://magi.nersc.gov, where users can upload their own 589 metabolite and FASTA files for analysis through MAGI.

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797 Figures and Tables

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800 Figure 1. MAGI workflow for consensus scoring. Mass spectrometry 801 features are connected to metabolites via methods such as accurate mass 802 searching or fragmentation pattern matching. These metabolites are expanded to include similar metabolites by using the Chemical Network. 803 These metabolites are then connected to reactions, which are reciprocally 804 805 linked to input gene sequences via homology (Reciprocal BLAST box). The 806 metabolite, reaction, and homology scores generated throughout the MAGI process are integrated to form MAGI scores (Scoring box). For details on MAGI 807 808 scores, see **Methods**.



809 810 Figure 2. MAGI associates more genes with reactions that can be ranked in S. coelicolor. a) Number of reactions associated with each gene 811 by MAGI, KEGG, and BioCyc. b) Venn diagram showing the genes connected 812 to one or more reactions by MAGI, KEGG, and/or BioCyc. c) Distributions of 813 the associations between a gene and a reaction for genes that have 814 815 annotations in MetaCyc or Kegg (orange), or are unique to MAGI (blue), highlighting that there are several high-scoring MAGI associations for genes 816 817 with no annotation. 818



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Figure 3. Pathway views of MAGI results. Metabolite, homology, and 820 integrative MAGI scores throughout the (a) actinorhodin and (b) menaguinone 821 822 biosynthesis pathways guides MAGI interpretations by visualizing results in a 823 broader context. Circular nodes represent metabolites, diamond nodes 824 represent reactions, and edges represent MAGI consensus scores. Border 825 color of circular nodes corresponds to the MIDAS metabolite score, and border width corresponds to the chemical network level searched in MAGI. Fill 826 827 color of diamond nodes correspond to the homology score. The line width of 828 the edges corresponds to the MAGI score. Abbreviations and legends for 829 metabolites and reactions are in supplementary table 8. The final step(s) in 830 the menaguinone biosynthesis are currently not known and are represented 831 by dashed edges and a "?" as the reaction.





834 Figure 4. Flowchart illustrating the key components of the MAGI 835 algorithm and process for associating undecylprodigiosin with **SC05896.** In the upper half of the flowchart, the mass spectrometry feature 836 837 with m/z 392.2720 at retention time 7.51 minutes was potentially identified 838 to be undecylprodigiosin, which is in the undecylprodigiosin synthase reaction. This reaction has no reference sequence, so could not be directly 839 840 connected to any S. coelicolor genes. Undecylprodigiosin was gueried for 841 similar metabolites in the chemical network, finding prodigiosin, which is in 842 the prodigiosin synthase reaction. This reaction does have a reference 843 sequence, which was used in a homology search against the S. coelicolor genome (Reaction to Gene BLAST), finding SCO5896 as the top hit. In the 844 845 lower half of the flowchart, the SCO5896 gene sequence was gueried against 846 the entire MAGI reaction reference sequence database in a homology search 847 (Gene to Reaction BLAST), finding the prodigiosin synthase and norprodigiosin synthase reactions. Norprodigiosin synthase did not have any 848 metabolomics evidence, The metabolite-to-reaction and gene-to-reaction 849 850 results were connected via the shared prodigiosin synthase reaction, effectively linking the feature 392.2720 to undecylprodigiosin and to 851 852 SCO5896.



854 855 Figure 5. Prioritization of MAGI gene function suggestions. a) Of the 1,883 MAGI-specific gene-metabolite linkages (Figure 2C), 591 genes were 856 associated with a reaction that was directly connected to an observed 857 metabolite (i.e. the chemical similarity network was not used to link a 858 859 metabolite to the reaction) (light blue). b) Of those, 190 genes had reciprocal agreement in bidirectional BLAST searches (light blue). c) Histogram of the 860 861 top MAGI scores of the 190 genes from panel (b). Through this process an actionable number of high-priority and high-strength novel gene function 862 863 hypotheses to test in future studies can be identified.

865Table 1. Comparison between MAGI, KEGG, and BioCyC annotations866for S. coelicolor genes discussed in this study.

| Gene | MAGI annotation (reaction) | MAG I scor e | Observed Metabolite Evidence | KEGG annotation (name) | KEGG Reaction Agreeme nt with MAGI | BioCyc annotati on (name) | BioCyc Reaction Agreeme nt with MAGI |
|---------|----------------------------------|-----------------------|---|---|--|------------------------------------|--|
| SCO4326 | RXN-10622 | 5.68 | Dihydroxy- naphthoate | 1,4-dihydroxy- 6-naphthoate synthase | Agree | ORF | None |
| SCO4327 | RHEA:25907 | 5.16 | Futalosine | None | None | ORF | None |
| SCO4494 | RXN-15264 | 5.57 | Carboxy- vinyloxy- benzoic acid | Aminodeoxy- futalosine synthase | Agree | ORF | None |
| SCO4506 | RXN-12345 | 5.57 | Carboxy- vinyloxy- benzoic acid | chorismate dehydratase | Agree | ORF | None |
| SCO4550 | RXN-10620 | 5.03 | Cyclic-DHFL | cyclic dehypoxanthi nyl futalosine synthase | Agree | ORF | None |
| SCO5074 | RXN1A0- 6312 | 5.37 | Bicyclic intermediate F & (S)- Hemiketal | None | None | ActVI- ORF3 | Agree |
| SCO5075 | RXN1A0- 6316 | 1.22 | Dihydro- kalafungin | None | None | ActVI- ORF4 | Agree |
| SCO5080 | RXN-18115 | 4.87 | DHK-red | 3-hydroxy- 9,10- secoandrosta- 1,3,5(10)- triene-9,17- dione monooxygena se [EC:1.14.14.1 2] | Disagree: R09819 | ActVA- ORF5 | Agree |
| SCO5081 | RXN1A0- 6318 | 4.63 | Dihydro- kalafungin | None | None | ActVA- ORF6 | Agree |
| SCO5091 | RXN1A0- 6307 | 5.95 | Bicyclic intermediate E | None | None | ActIV | Agree |
| SC05315 | RXN-15413 | 4.58 | WhiE_20C_su bstrate | None | None | Polyketide aromatas e | None |

| SCO5896 | RXN-15787* | 1.32 | Undecyl- prodigiosin | pyruvate, water dikinase | Disagree: R00199 | RedH | Agree* |
|---------|------------|------|-------------------------|---|---|-----------|--------|
| SCO6300 | RXN0-5226 | 3.22 | Anhydro-NAM | beta-N-acetyl- hexosaminida se | Disagree: R00022, R05963, R07809, R07810, R10831 | hydrolase | None |
| SCO7595 | RHEA:24952 | 5.23 | Anhydro-NAM | anhydro-N- acetylmurami c acid kinase | None | ORF | None |

* Due to chemical network search, this reaction was listed as the prodigiosin 867

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synthase reaction but the metabolite connected to it was undecylprodigiosin, requiring manual interpretation to determine the actual reaction connected 869

to the gene was undecylprodigiosin synthase. 870