Lawrence Berkeley National Laboratory

Biological Systems & Engineering

Title

PeakDecoder enables machine learning-based metabolite annotation and accurate profiling in multidimensional mass spectrometry measurements

Permalink

https://escholarship.org/uc/item/6mx6n2bc

Journal Nature Communications, 14(1)

ISSN 2041-1723

Authors

Bilbao, Aivett Munoz, Nathalie Kim, Joonhoon <u>et al.</u>

Publication Date

2023

DOI

10.1038/s41467-023-37031-9

Copyright Information

This work is made available under the terms of a Creative Commons Attribution-ShareAlike License, available at https://creativecommons.org/licenses/by-sa/4.0/

Peer reviewed

1 PeakDecoder enables machine learning-based metabolite annotation

2 and accurate profiling in multidimensional mass spectrometry

3 measurements

4

Aivett Bilbao^{1,2*}, Nathalie Munoz^{1,2}, Joonhoon Kim^{1,2}, Daniel J Orton¹, Yuqian Gao^{1,2}, Kunal
Poorey³, Kyle R. Pomraning^{1,2}, Karl Weitz¹, Meagan Burnet¹, Carrie D. Nicora¹, Rosemarie
Wilton^{4,2}, Shuang Deng^{1,2}, Ziyu Dai^{1,2}, Ethan Oksen⁵, Aaron Gee⁶, Rick A. Fasani⁶, Anya
Tsalenko⁶, Deepti Tanjore^{5,2}, James Gardner^{5,2}, Richard D. Smith¹, Joshua K. Michener^{7,2}, John
M. Gladden^{3,2}, Erin S. Baker⁸, Christopher J. Petzold^{5,2}, Young-Mo Kim^{1,2}, Alex Apffel⁶, Jon K.

- 10 Magnuson^{1,2} and Kristin E. Burnum-Johnson^{1,2*}
- 11
- 12¹ Pacific Northwest National Laboratory, Richland, WA, USA
- 13 ² US Department of Energy, Agile BioFoundry, Emeryville, CA, USA
- 14 ³ Sandia National Laboratory, Livermore, CA, USA
- 15 ⁴ Argonne National Laboratory, Lemont, IL, USA
- 16 ⁵ Lawrence Berkeley National Laboratory, Berkeley, CA, USA
- 17 ⁶ Agilent Research Laboratories, Agilent Technologies, Santa Clara, CA, USA
- 18 ⁷ Oak Ridge National Laboratory, Oak Ridge, TN, USA
- ⁸ Department of Chemistry, University of North Carolina, Chapel Hill, NC, USA
- 20
- 21 These authors contributed equally: Aivett Bilbao, Nathalie Munoz, and Joonhoon Kim.
- 22

23 * Correspondence:

24 Aivett.Bilbao@pnnl.gov and Kristin.Burnum-Johnson@pnnl.gov

25 Abstract

26 Multidimensional measurements using state-of-the-art separations and mass spectrometry 27 provide advantages in untargeted metabolomics analyses for studying biological and 28 environmental bio-chemical processes. However, the lack of rapid analytical methods and robust 29 algorithms for these heterogeneous data has limited its application. Here, we develop and 30 evaluate a sensitive and high-throughput analytical and computational workflow to enable 31 accurate metabolite profiling. Our workflow combines liquid chromatography, ion mobility 32 spectrometry and data-independent acquisition mass spectrometry with PeakDecoder, a machine 33 learning-based algorithm that learns to distinguish true co-elution and co-mobility from raw data 34 and calculates metabolite identification error rates. We apply PeakDecoder for metabolite profiling 35 of various engineered strains of Aspergillus pseudoterreus, Aspergillus niger, Pseudomonas 36 putida and Rhodosporidium toruloides. Results, validated manually and against selected reaction monitoring and gas-chromatography platforms, show that 2683 features could be confidently 37 38 annotated and quantified across 116 microbial sample runs using a library built from 64 standards. 39

40 Introduction

41 Metabolomics is the study of the small molecules produced by complex networks of cellular 42 processes and biochemical reactions in living systems. Metabolites are the end point of the flow 43 of information from DNA to the biological phenotype and represent chemical fingerprints directly 44 reflecting the physiological conditions, intracellular regulation, and effects that environmental 45 factors induce in biological cells or organisms. As such, metabolomics helps in a variety of 46 applications, from understanding disease progression in clinical settings to estimating 47 overproduction for metabolic engineering^{1,2}.

48

Advances in synthetic biology, genome editing, and DNA synthesis capabilities have propelled the ability to routinely design and generate thousands of novel strains for biomanufacturing research. The Agile BioFoundry (ABF) consortium of national laboratories utilizes state-of-the-art capabilities within the framework of the Design, Build, Test, and Learn (DBTL) cycle to develop engineered organisms³. Accurate analytical tools with fast turnaround time in Test are critical in developing microorganisms that can produce desired fuels and chemicals from renewable biological feedstocks.

56

57 The most popular and widely used analytical platform for the analysis of metabolic species in 58 complex mixtures is mass spectrometry (MS) combined with liquid chromatography (LC) or gas 59 chromatography (GC) separations^{2,4,5}. However, hundreds to thousands of primary and 60 secondary metabolites in nature display a high degree of structural diversity with many isomers 61 and nominal mass isobars that co-elute and have similar fragmentation patterns, all of which 62 constitute a significant analytical challenge in terms of detection and annotation. The incorporation 63 of several orthogonal technologies in MS-based workflows can provide heterogeneous 64 information to tackle these challenges. In fact, experimental measures such as retention time (RT) 65 from chromatography, collision cross-section (CCS) from ion mobility spectrometry (IM) or stable 66 isotope labeling, are necessary to complement MS/MS similarity and add confidence in overall 67 compound identification workflows⁶.

68

69 Besides increasing annotation confidence, multidimensional LC-IM-MS workflows collecting 70 extensive fragmentation spectra with data-independent acquisition (DIA) methods are providing 71 heterogeneous information which allows deeper understanding in metabolomics studies. IM is a 72 gas phase separation technique increasingly used to distinguish structurally similar molecules, 73 isomers and molecular classes in biological and environmental samples⁷. Unlike LC that 74 separates molecules based on hydrophobicity, IM separates gas-phase molecular ions based on 75 their charge, size, and shape, which improves selectivity and coverage compared to routine LC-76 MS-based methods.

77

In DIA the mass spectrometer is operated to systematically collect multiplexed fragment-ion spectra (MS2) from all detectable precursors (MS1) within a wide *m/z* range and in a single chromatographic run, independently of their intensities⁸. Like initially found in proteomics⁹, in metabolomics the MS2 spectrum quality of ions that get selected during standard data-dependent acquisition (DDA) is higher, but the overall MS2 coverage and quantitative precision using DIA is 83 better¹⁰. While DIA provides increased reproducibility and guantitation performance, it requires 84 more elaborated processing algorithms compared to DDA. Two main DIA processing strategies initially established for proteomics have been adapted to metabolomics in a handful of DIA 85 metabolomics tools. The first strategy applies untargeted feature detection followed by 86 87 deconvolution of fragment ion spectra (here referred to as UFD). A popular tool used for UFD in metabolomics is MS-DIAL¹¹, which groups precursors and their corresponding fragments based 88 89 on the similarity of their elution profiles, generates pseudo-MS2 spectra and matches them 90 against a reference MS2 library. Other reported tools applying UFD are MetaboDIA¹² and 91 DaDIA¹³. The second DIA algorithmic strategy employs targeted data extraction (here referred to 92 as TDX). TDX requires a library of target analytes with retention times, and precursors with 93 corresponding fragment masses, which are utilized as coordinates to mine the DIA spectra and 94 generate extracted ion chromatograms (XIC) for precursor and fragments per target analyte, as 95 the so-called 'peak-group'. Multiple sub-scores are then calculated per peak-group to assess 96 coelution and identification. Software employing TDX include Skyline¹⁴, MetDIA¹⁵ and 97 DIAMetAlyzer¹⁶. Another tool demonstrated for DIA using a different approach is DecoID¹⁷, where 98 the MS2 deconvolution is achieved by mixing database spectra to match an experimentally 99 acquired spectrum using least absolute shrinkage and selection operator (LASSO) regression.

100

101 While these tools exist for DIA metabolomics, new tools capable to fully exploit all dimensions 102 with controlled error rates in multidimensional LC-IM-MS measurements with DIA spectra are 103 needed. Skyline and MS-DIAL were adapted to support the additional IM dimension but they do 104 not provide a false-discovery rate (FDR) control method. Unlike proteomics, the field of 105 metabolomics still lacks a generally accepted, validated, and automated calculation of error rates 106 for MS2 compound identification with FDR assessments¹⁸. Several methods have been proposed 107 to generate decoys and estimate FDR in metabolomics. For imaging-MS, pySM¹⁹ generates 108 decoys by using implausible ion adducts. For DDA, Passatutto²⁰ uses re-rooted fragmentation 109 trees, JUMPm²¹ adds a small odd numbers of hydrogen atoms, and XY-Meta²² combines original 110 and randomly selected MS2 peaks. And recently reported for DIA, DIAMetAlyzer¹⁶, provides an 111 FDR estimation employing Passatutto²⁰ but it does not support the IM separation. These methods rely on annotated spectra or a sample-specific metabolite database for FDR estimation. 112

113

114 Here, we develop a sensitive and high-throughput analytical and computational workflow that 115 combines LC-IM-MS multidimensional measurements with PeakDecoder, an algorithm that automatically calculates error rates for metabolite identification, independently of spectral 116 117 annotations or libraries. PeakDecoder proposes an alternative method for decoy generation from 118 raw DIA spectra, incorporating concepts from DIA and spectral library searching into a machine 119 learning (ML) strategy that combines both UFD and TDX. To illustrate our metabolomics workflow 120 and demonstrate its utility, we apply it to study microbial samples from various strains engineered 121 under projects of the ABF consortium.

122 **Results**

123 Optimizing the LC-IM-MS analytical method

124 We defined a list of 64 metabolites of interest for the study of various strains of *Pseudomonas* 125 putida, Aspergillus pseudoterreus, Aspergillus niger and Rhodosporidium toruloides, all relevant 126 microorganisms in the biotechnology field for production of value-added chemicals. The panel 127 consisted of metabolites from central carbon metabolism including glycolysis, tricarboxylic acid 128 cycle (TCA) cycle intermediates, amino acids, and 'coenzyme A' molecules (CoAs) that are 129 routinely analyzed in ABF studies to obtain an overview of changes in the metabolism of cells. 130 Additionally, metabolites specific to ABF host-bioproduct pairs, meaning compounds that are 131 directly along the engineered pathways were also included.

132

133 The microorganisms in this study are promising industrial hosts and have a variety of application interests. P. putida is a Gram-negative, rod-shaped bacterium that is metabolically versatile, 134 135 tolerant to toxins and solvents, with a high supply of reducing power, making it ideal for numerous 136 biomanufacturing applications²³. The eukaryotic microorganisms A. pseudoterreus and A. niger 137 (filamentous fungi) were modified for production of 3-hydroxypropionic acid, a polymer precursor 138 that can be dehydrated to produce acrylic acid and can be used directly within existing 139 infrastructure²⁴. Similarly, the *R. toruloides* (oleaginous yeast) strains were engineered for 140 production of bisabolene, which is a precursor to a diesel alternative and is considered an ideal 141 platform for bioconversion of lignocellulose into lipids and related chemicals²⁵.

142

143 To select an LC method, we implemented the Automated chromatographic Method Selection 144 Software (AMSS), which utilizes chemical and physical properties of metabolites to predict the LC 145 method that maximizes the number of metabolites detected (see Methods). The evaluation of the 146 selected metabolites using AMSS predicted HILIC with negative ESI as the best method 147 (Supplementary Fig. 1). The LC conditions were first implemented and optimized by selected 148 reaction monitoring (SRM) analyses of a subset of the standards and led to a total acquisition 149 time of 9 min per run. Compared to the methods typically used to perform GC-MS-based global 150 metabolomics²⁴, this LC method provides a \sim 3x faster sample analysis time and can detect other 151 molecules which are undetectable by GC such as CoAs. DDA methods with short LC separation 152 (<15 min) would be limited to only select the top 3-5 ions^{18,26} per cycle to preserve the MS1 153 sampling rate and quantitation dynamic range, which in turn would result in MS/MS under 154 sampling of medium-low-abundance ions. Therefore, after initial optimization the same LC system 155 was utilized to perform the LC-IM-MS analyses in the All-Ions DIA mode (Fig. 1). A library with 156 RT, CCS and transitions (hereafter referred to as precursor and fragments) was built from the 157 analysis of standards in deprotonated ion form. The list of metabolites can be found in 158 Supplementary Table 1 and the library can be found in Supplementary Data 1. To evaluate the 159 LC-IM-MS system against the gold standard SRM platform, dilution experiments were performed 160 using representative standards. Supplementary Fig. 2 shows the calibration curves with linearity 161 and increased sensitivity of LC-IM-MS over SRM for concentrations to as low as 0.075 pmol and 162 covering four orders of magnitude. Next, 81 microbial samples from the various ABF-engineered 163 hosts and conditions were analyzed by LC-IM-MS and processed using PeakDecoder.

164 Developing the PeakDecoder algorithm

We implemented an alternative scoring algorithm for DIA metabolite identification which uses a 165 166 'raw spectrum centric' approach with UFD for ML training and a 'metabolite centric' approach with 167 TDX for metabolite scoring (i.e., ML inference). Using only the unannotated LC-IM-MS DIA 168 experimental spectra from biological samples, PeakDecoder learns to discriminate true co-elution 169 (and co-mobility) of a precursor and its fragments from poor co-elution undistinguishable from 170 random chance. As Fig. 2-a shows, the PeakDecoder workflow has six steps for ML training and 171 inference. First, the LC-IM-MS DIA data from the biological samples is processed in UFD mode 172 using MS-DIAL¹¹. Second, a preliminary training set is generated by using the detected and 173 deconvoluted peak-groups as targets and producing their corresponding decoys. Third, TDX is 174 performed using Skyline¹⁴ to extract the precursor and fragment ion signals for the training set 175 from all the LC-IM-MS DIA runs and export their XIC metrics. Fourth, a final training set is 176 generated applying filtering for high-quality fragments to keep high-quality peak-groups as targets 177 (i.e., precursor S/N > 20, and at least 2 fragments with mass error < 15 ppm, RT difference to 178 their precursor < 0.1 min, and FWHM difference to their precursor larger than 2x the precursor 179 FWHM; details in Methods) and their corresponding decoys. A support vector machine (SVM) 180 classifier is trained using multiple scores calculated from the XIC metrics of each peak-group in 181 the training set: the cosine similarity between the expected and XIC intensities, and the mean and 182 standard deviation of each precursor and its fragments for RT, LC-FWHM and mass error metrics 183 (details in Methods). These scores are used as ML features which measure co-elution and 184 similarity to the expected values²⁷. After scoring the training set, the true and false positives can 185 be used to estimate an FDR. Fifth, TDX is performed to extract the signals of the query set of 186 metabolites in the library from all the LC-IM-MS runs and export their XIC metrics. Finally, the 187 trained model is used to score the query set of metabolites and results can be filtered using the 188 PeakDecoder score corresponding to the estimated FDR threshold from a table with pairs of 189 values (FDR, PeakDecoder score) automatically generated.

190

191 PeakDecoder takes advantage of DIA spectra, where the combination of precursor and its 192 fragments enable selective and sensitive detection of a molecule by a peak-group of co-eluting 193 fragment ion chromatograms²⁸. Our algorithm is similar to the mProphet scoring method in terms 194 of using decoy transitions^{29,30}. The mProphet method introduced the concept of decoy transitions 195 at the measurement level for SRM proteomics, and it was later adapted at the data extraction 196 level for DIA. The decoy transitions are used to optimize a combination of the available individual 197 scores and to derive statistical error rate estimates by parameterizing a null distribution. However, 198 decoys in those original methods are generated from the protein database by reversing or 199 shuffling the sequences. Due to the much larger structural diversity, more complex fragmentation 200 mechanism and ubiquitous isomers compared to peptides, such decoy generation methods 201 cannot be applied for small molecules. In contrast, PeakDecoder generates the decoys from the 202 high-quality peak-groups deconvoluted from the LC-IM-MS DIA experimental spectra of the 203 biological samples.

204

Methods to generate decoys from experimental spectra have been previously reported, however, from a DDA MS/MS target library (i.e. annotated spectra), first in proteomics^{31,32} and more recently 207 in metabolomics^{20,22}. We propose an alternative strategy to generate decoys taking advantage of 208 the comprehensive nature of the DIA spectra. Instead of generating decoys from the target library, 209 we perform UFD and TDX in the LC-IM-MS DIA data to generate a training set of peak-groups. 210 The high-quality peak-groups constituted by the detected precursors (MS1) and its deconvoluted 211 fragments (i.e., pseudo MS2) are used as targets. This strategy provides a noise-filtered 'clean' 212 set of targets which was reported to be necessary to reach accurate estimates in spectrum level 213 decoy-based methods²⁰. We then employ a pairing and swapping strategy, similar to the 214 precursor-swap method proposed by Cheng et. al.³², but rather than swapping precursors, we 215 generate the respective decoy precursors and fragments from the same targets by swapping pairs 216 of fragment m/z (Fig. 2-b). Pairing precursors with the same number of fragments was used as 217 an approach to increase the chances that the molecules are similar and to ensure that the overall 218 distributions of general properties of targets and decoys are the same. Generating decoys by 219 pooling and randomly adding fragments was avoided because it has previously shown poor 220 performance (naïve method)²⁰, as it increases the probability of generating unrealistic decoys. 221 Since the deconvoluted data represent real molecules, our decoy strategy is valid in practice and 222 the generated decoys comply with several conditions or properties, previously proposed for 223 proteomics³¹, to calculate FDR with a valid target-decoy model: (i) the decoy library has the same 224 precursor m/z and charge distributions as the target library. (ii) target and decoy spectra include 225 the same number of peaks and have the same intensity sum distribution, and (iii) decoy spectrum 226 peaks are positioned on realistic m/z values (fragments that naturally occur).

227

228 Contrary to previously proposed methods for FDR assessment that rely on large libraries of 229 annotated MS/MS spectra, PeakDecoder was designed to confidently identify metabolites from 230 libraries, but independently of the number of metabolites in the library. The estimated error rates 231 are independent of any library and therefore experimental or in-silico generated libraries of any 232 size could be potentially utilized. The scoring becomes 'metabolite centric' and provides the 233 probability that a given metabolite is present in the sample based on the quality of its detected 234 signals in the LC-IM-MS DIA data. After the model is trained directly from the unannotated LC-235 IM-MS DIA data, it can be used to automatically score metabolites in libraries.

236

237 Since PeakDecoder generates the decoys from unannotated LC-IM-MS DIA experimental 238 spectra, the size of the target library does not affect its performance. However, the performance 239 of PeakDecoder depends on the training set and the validity of the estimated FDR depends on 240 the number of generated false positives. The size and quality of the training set can be controlled 241 in two ways: the parameters of the UFD tool used to generate the preliminary training set (Fig. 2-242 a, Step-1) and the filtering for high-quality fragments used to generate the final training set (Fig. 243 2-a, Step-4). At the same time, a tradeoff in the quality of peak-groups is necessary to avoid 244 overfitting and perfect training accuracy, and thus, to estimate a reliable FDR. These components 245 allow the user to define the quality of the resulting annotations and are evaluated using microbial 246 data in the next section.

247 Applying PeakDecoder in microbial samples

248 We processed the microbial LC-IM-MS data using PeakDecoder. The datasets represented 249 varied sample complexity and feature density: low for A. pseudoterreus & A. niger, medium for P. 250 putida and high for *R. toruloides*. Supplementary Fig. 3 shows the distributions of ions illustrating 251 the general properties of the targets and decoys generated for training. Fig. 3 shows results for 252 the *P. putida* samples. The PeakDecoder score which combines individual scores provided an 253 improved discrimination power between targets and decoys (Fig. 3-a). An example of 254 chromatograms and filtered IM window for 'fructose 1,6-diphosphate (F16DP)' from the standard 255 (precursor m/z 338.98877, RT 4.95 min, CCS 155.00 and 6 fragments) and a microbial sample is 256 shown in Fig. 3-b, confidently identified with a PeakDecoder score of 0.9966 and 0.005 q-value. 257 Supplementary Fig. 4 shows the PeakDecoder training performance for all microbial samples and 258 a summary is shown in Table 1. A total of 2683 features could be confidently annotated. 259 Annotations could be attributed to either all dimensions by RT-CCS-DIA or to RT-CCS for features 260 without detected fragments (i.e., MS1 level only). The number of features annotated in each 261 dataset includes replicates and is independent of the number of unique metabolites identified. For 262 instance, in the case of the A. pseudoterreus & A. niger dataset, many more features were 263 annotated, indicating that metabolites were detected in multiple replicates across all sample 264 conditions.

265

266 To control the size and quality of the final training set, we defined the parameters of the UFD tool 267 (Fig. 2-a, Step-1) and the filtering for high-quality fragments (Fig. 2-a, Step-4) according to the 268 characteristics of our analytical method and instrumentation (e.g., fragments with RT difference 269 to their precursor < 0.1 min) and annotation guality preferred (e.g., at least 3 fragments). Because 270 of the low sample complexity of the A. pseudoterreus & A. niger dataset, a smaller number of 271 deconvoluted peak-groups were detected, therefore only 234 target peaks could be generated for 272 training and were not sufficient for a good FDR estimation. The medium sample complexity of the 273 P. putida dataset provided the best FDR estimation. Supplementary Fig. 5 shows that the training 274 performance was not significantly impacted by the deconvolution parameters if the numbers of 275 targets was sufficient (accuracy > 98.86 if the resulting training set contained between 2760 and 276 6720 targets), but at the same time, if the classifier resulted in a close-to-perfect accuracy (>99), 277 the minimum non-zero FDR that could be estimated was affected due to the small number of false 278 positives. Conversely, the high sample complexity in the R. toruloides dataset resulted in poor 279 performance when using the default filtering for high-quality fragments generating 8674 280 targets/decoys for training, where the minimum non-zero estimated FDR for the highest 281 PeakDecoder score was 3% (Supplementary Fig. 6-a). Stringent values were applied to filter the 282 high-quality fragments generating 1400 targets/decoys (Supplementary Fig. 6-b) and a minimum 283 non-zero estimated FDR of 1% could be obtained. The results indicate that more training data 284 does not translate into higher accuracy and further improvements for filtering high-quality 285 fragments (i.e., generating a smaller training set with peak-groups of appropriate quality) could 286 be needed for datasets with high sample complexity. Supplementary Fig. 6-c depicts results from 287 a deuterated standard (tryptophan d5) spiked in solvents and in a microbial sample matrix. 288

289 In all samples, PeakDecoder could identify the metabolites expected in at least one condition of 290 each microbial dataset (a list of unique metabolites generated by manual inspection of the most 291 intense replicates). A handful of cases that were missed initially were recovered after manual 292 correction of the Skyline chromatogram peak detection. Supplementary Tables 2-4 show the 293 scores and annotation confidence level (best replicate per metabolite). In addition, we performed 294 targeted analyses of a subset of metabolites by SRM in P. putida samples and GC-MS analyses 295 of A. pseudoterreus and A. niger samples to further evaluate the performance of our method in 296 biological samples (Supplementary Fig. 7). Similar trends were observed for the metabolites 297 identified in common by the different platforms.

298

299 Comparing PeakDecoder to other workflows

300 UFD (MS-DIAL) and TDX (Skyline) are two different approaches with different advantages and 301 disadvantages. While the UFD does not rely on a library and high-quality peak-groups from its 302 deconvolution results can be used for training, applying TDX offers advantages over UFD for 303 annotation in DIA, particularly for All-lons data, where the full mass range is co-fragmented, and 304 the likelihood of interference greatly increases as sample complexity increases. In complex 305 samples, multiple precursors with very similar RT and DT are present as a series of partial 306 overlapping ions which compromise the effectiveness of UFD algorithms. However, when 307 performing TDX, only the relevant chromatograms are extracted in a directed and highly selective 308 fashion.

309

310 PeakDecoder combines both UFD and TDX strategies and addresses limitations in the respective 311 existing tools. Specifically, the re-extraction of signals by TDX in Skyline allows specifying a DT 312 offset for fragments characteristic for the IM instrumentation³³ (see Methods), which is not applied 313 in MS-DIAL and results in a poor deconvolution of fragments with the smaller masses. On the 314 other hand, the UFD in MS-DIAL allows accurate CCS evaluation using the experimental CCS 315 values, which is not available in Skyline because it does not perform a peak detection in the IM 316 dimension and is limited to use the CCS information as a filtering window (e.g., Fig. 3-b). Besides 317 combining the best features of these two tools, PeakDecoder uses the peak shape metrics, 318 combines the individual scores into a composite score, and allows FDR estimation, all of which 319 are impossible with MS-DIAL or Skyline alone.

320

321 To benchmark PeakDecoder and evaluate the reliability of our FDR estimation, we performed a 322 comparison against the ground truth generated from manually curating the full P. putida dataset, 323 with 550 peak-groups including 233 positives and 317 negatives (Fig. 3-c). Due to the poor 324 deconvolution of fragments with the smaller masses, MS-DIAL resulted in the lowest number of 325 true positives (TP=70), even when using a relaxed threshold for its total score (> 60). While 326 PeakDecoder at 1% estimated FDR missed 4 TP compared to Skyline (cosine similarity > 0.8), it 327 decreased the number of false positive annotations (FP: PeakDecoder=4, MS-DIAL=13, 328 Skyline=15). The estimated 1% FDR corresponded to a $\sim 2\%$ actual FDR, and while there is an 329 underestimation and the results are limited by our small library, they show that PeakDecoder 330 could be used to filter out FP.

331

332 Our decoy strategy for DIA data together with IM and LC conveys a powerful multidimensional 333 characterization of metabolites that address several important challenges. For many metabolites 334 only a few characteristic fragment ions can be detected, rendering the use of classic spectral 335 similarity searches unreliable¹⁸. Moreover, some metabolites could not be detected with even a 336 single fragment. In these cases, the CCS increased the identification confidence compared to 337 using the RT and accurate mass alone. Since our library was built from pure standards, even for 338 standards without fragments, the identification based on RT and CCS could be considered as a 339 confidence of "Level 1" according to the Metabolomics Standards Initiative³⁴, as they are two 340 different analytical techniques. Besides, multidimensional LC-IM-MS increases the separation, 341 important for metabolites that co-elute, where DIA alone is challenged by fragments common to 342 co-eluting metabolites. Fig. 4 illustrates the power of multidimensional separations to increase the 343 selectivity and therefore increase the annotation confidence and quantitation accuracy. The 344 number of possible LC-IM-MS peaks from MS-DIAL untargeted feature detection results matched 345 within tolerances (0.01 mass, 0.2 min RT, and 0.8% CCS) was reduced when using all 346 dimensions. High IM resolving power is essential for small molecules and current IM instruments 347 are able to separate CCS differences as low as 0.8%.

348 Metabolomics of *A. pseudoterreus and A. niger* strains

349 PeakDecoder was applied for metabolomics profiling of A. pseudoterreus and A. niger strains 350 engineered to produce 3-hydroxypropanoic acid (3HP) using the β -alanine pathway³⁵. Our three 351 engineered A. pseudoterreus strains²⁴ with varying levels of 3HP production (low, medium, and 352 high) and their parent strain (ATCC 32359 ∆cad: cis-aconitic acid decarboxylase deletion) were 353 analyzed. Since the engineered A. pseudoterreus strains produced significant amount of other 354 organic acids²⁴, we also developed and profiled A. niger strains engineered with the same β -355 alanine pathway. Five engineered A. niger strains exhibiting different levels of 3HP production 356 (low, medium, high, higher, and highest) and their parent strain (ATCC 11414) were included.

357

358 Metabolomics profiling of 3HP-producing *A. pseudoterreus* and *A. niger* strains revealed species-359 specific metabolic responses to increasing 3HP production. Specifically, we found that L-360 aspartate, the precursor to the β -alanine 3HP production pathway, showed very little change in 361 3HP producing *A. pseudoterreus* strains, while its level decreased significantly in 3HP producing 362 *A. niger* strains (Fig. 5). In the β -alanine 3HP production pathway, L-aspartate is converted to 363 3HP via β -alanine using multiple aminotransferases.

- 364
- 365 L-aspartate -> β -alanine + CO₂
- 366 β -alanine + pyruvate -> malonate semialdehyde + L-alanine
- 367 L-alanine + α -ketoglutarate -> pyruvate + L-glutamate
- 368 oxaloacetate + L-glutamate -> L-aspartate + α -ketoglutarate
- 369 (net reaction) β -alanine + oxaloacetate -> malonate semialdehyde + L-aspartate
- 370 malonate semialdehyde + NADPH + H⁺ -> 3HP + NADP⁺

371

372 L-aspartate is first converted to β -alanine by aspartate-1-decarboxylase, and the amino group 373 from β -alanine is transferred to pyruvate yielding malonate semialdehyde and L-alanine by β -374 alanine/pyruvate aminotransferase. Malonate semialdehyde is converted to the final product 3HP 375 by 3-hydroxypropionate dehydrogenase, but L-alanine needs be converted back to pyruvate by 376 alanine transaminase by transferring the amino group to α -ketoglutarate generating L-glutamate. 377 The aminotransferase cycle can be closed by generating the precursor L-aspartate by transferring 378 the amino group from L-glutamate to oxaloacetate. Therefore, the amino group acceptor and 379 donor pairs (pyruvate/L-alanine and α -ketoglutarate/L-glutamate) play an important role in the β -380 alanine 3HP production pathway. 381

382 PeakDecoder allowed us to investigate the changes in these amino group acceptor and donor 383 pairs as well as undesired byproducts such as 4-aminobutyric acid and 2,4-aminobutanoic acid. 384 Similar to the conversion of L-aspartate to 3HP via β -alanine, L-glutamate can be converted to 385 succinate via 4-aminobutyric acid (GABA). In the GABA degradation pathway, GABA is first 386 deaminated to succinate semialdehyde by 4-aminobutyrate aminotransferase UGA1 using α -387 ketoglutarate/L-glutamate pair, which was one of the most significantly upregulated enzymes in 388 the engineered A. pseudoterreus strains in our previous study²⁴. In this study, we observed 389 significantly decreased levels of succinate semialdehyde in A. niger strains producing high levels 390 of 3HP using the developed workflow, confirming that the engineered 3HP pathway is affecting 391 the GABA degradation pathway in A. niger as well. We also previously hypothesized that the 392 promiscuous activity of upregulated UGA1 resulted in the accumulation of 2,4-diaminobutyric acid 393 from L-aspartate via L-aspartate 4-semialdehyde in A. pseudoterreus. Here, we found that the 394 accumulation of 2,4-diaminobutyric acid was not consistently observed in the engineered A. niger 395 strains in contrast to the observation in A. pseudoterreus. This is likely due to the significantly 396 decreased level of the precursor L-aspartate in the engineered A. niger strains. The level of L-397 aspartate 4-semialdehyde is consistently lower in the engineered A. pseudoterreus strains, but 398 not in the engineered A. niger strains.

399 Omics of engineered muconate-catabolizing *P. putida* strains

400 P. putida has biochemical properties that make it ideal for hosting biochemical transformations³⁶. 401 Due to its naturally diverse and flexible catabolism it can metabolize aliphatic, aromatic, and 402 heterocyclic compounds in addition to glucose³⁷. To use *P. putida* for industrial bioprocessing, 403 genetic modifications must be incorporated into the strains requiring the expression of 404 heterologous genes and pathways. Chaves et al. studied the importance of chromosomal 405 integration location, which affects heterologous protein expression independent of typical design parameters such as copy number, promoter, and terminator type³⁷. Wild-type (WT) P. putida 406 407 KT2440 cannot grow on cis, cis-muconate as a sole carbon source, despite using this compound 408 as a key intermediate in aromatic catabolism. To enable muconate catabolism, a transmembrane 409 transporter for muconate (mucK) was integrated into three different chromosomal sites (PP2224, 410 PP1642, and PP5042). Samples with the transmembrane transporter were grown in M9 minimal 411 medium supplemented with 30 mM muconate and analyzed using a targeted proteomics 412 approach to quantify the amount of muck present. The previous results showed that the growth 413 rate with muconate inversely correlated with the expression levels of the transporter.

414

415 To provide additional insights into the metabolic changes during growth with a new carbon 416 substrate, we used PeakDecoder to quantify metabolites in *P. putida* WT grown on glucose and 417 mucK (PP2224, PP1642 and PP5042) grown on muconate. Compared to WT, muconate-418 catabolizing *P. putida* mucK strains showed decreased levels of metabolites in the ED-EMP cycle 419 (glucose utilization) such as fructose 6-phosphate, fructose 1,6-diphosphate and glyceraldehyde 420 3-phosphate among others (Fig. 6). Targeted proteomics, performed on the same cell pellets of 421 muck strains, showed a corresponding decrease in enzymes that are part of ED-EMP pathway 422 and in levels of pyruvate dehydrogenase and pyruvate carboxylase, which catalyze the 423 conversion of pyruvate to acetyl-CoA and to oxaloacetate, respectively. These lower levels match 424 with the accumulation of pyruvate in mucK strains cultured with muconate. Accumulation of 425 pyruvate had also been observed in *P. putida* grown in a glucose:benzoate mixture vs glucose 426 alone³⁸. In contrast, increased levels of metabolites (α -ketoglutarate, fumarate, malate) and 427 enzymes from the TCA cycle were observed in the mucK strains. Levels of enzymes at the 428 entrance point of acetyl-CoA into TCA cycle and those routing succinyl-CoA and succinate into 429 TCA cycle were upregulated in mucK compared to WT. Muconate is metabolized via the β-430 ketoadipate pathway before joining the central carbon metabolism via acetyl-CoA and succinate. 431 Although no metabolites in the beta-ketoadipate pathway were detected, enzymes in this pathway 432 were significantly upregulated in the mucK strains compared to WT which is expected considering 433 muconate was used as the carbon source. Changes in metabolite levels in peripheral pathways 434 were also clear. Gluconate and 2-ketogluconate were lower or not detected in mucK compared 435 to WT which is in line with the absence of glucose supplementation in the strains with the 436 transporter. These results suggest a shift in metabolism supported on succinate and acetyl-CoA 437 fueling the TCA cycle from the β -ketoadipate pathway and less reliance on ED-EMP pathway 438 when muconate is used as carbon source. Similar results were observed when P. putida was 439 grown in p-coumarate³⁹. Supplementary Fig. 8 shows the targeted proteomics guantitation results and changes in metabolic pathways for mucK PP1642 and mucK PP2224 compared to the WT. 440

441 Mevalonate pathway in *R. toruloides* strains

442 R. toruloides is an important model microorganism for synthetic biology and industrial 443 biotechnology due to its capacity to bioconvert lignin, the most underutilized component of plant 444 biomass⁴⁰. Metabolic engineering of *R. toruloides* can generate distinct bio-products including 445 bisabolene, the immediate precursor of bisabolane and an alternative to D2 diesel fuel⁴¹. For 446 example, the Agile BioFoundry R. toruloides strain, GB2, can produce bisabolene in high 447 quantities of 2.2 g/L from lignocellulosic hydrolysate in 2-L fermenters⁴². Another key advantage 448 of R. toruloides is that it can grow on mixed-carbon sources and tolerate growth inhibitors often 449 present in lignocellulosic hydrolysates⁴⁰. However, these hydrolysates present a significant 450 challenge in biochemical conversion due to feedstock variability⁴³.

451

We employed PeakDecoder and global proteomics analyses to characterize *R. toruloides* GB2 cultured on lignocellulosic hydrolysates derived from corn stover with variable levels of ash (A) and moisture (M), each parameter cataloged as High (H) or Low (L) and the 4 possible combinations of them (HAHM, HALM, LAHM, and LALM). Samples were collected at two time points of fermentation, during exponential growth (36 hr) and at the end of this growth phase (60
hr). A total of 37 unique metabolites were confidently detected in at least one sample and
quantified across all samples (Supplementary Fig. 9).

459

460 Bisabolene is produced upon the introduction of bisabolene synthase and its precursor, farnesyl 461 pyrophosphate (FPP), is part of the mevalonate pathway. Fig. 7 details our mevalonate pathway 462 metabolomic and proteomic results. The levels of HMG-CoA were significantly higher in cells 463 cultured in high ash low moisture (HALM) conditions at 60 hours. The rate limiting step in the 464 mevalonate pathway is the conversion of HMG-CoA to mevalonic acid by 3-hydroxy-3-465 methylglutaryl-coenzyme A reductase (HMGR)⁴⁴. R. toruloides, like mammalian systems, has 466 only one HMGR gene⁴⁵ and mammalian HMGR and yeast Hmg2p (from Saccharomyces 467 *cerevisiae*) are both subject to feedback control by the sterol pathway⁴⁶. Previous studies 468 identified FPP or FPP derivatives as the positive signal for HMGR degradation in yeast^{44,46}. Here 469 we detected isopentenyl pyrophosphate/dimethylallyl pyrophosphate (IPP/DMAPP) and geranyl 470 pyrophosphate GPP. IPP and DMAPP are isomeric molecules which could not be separated by 471 the current drift tube IM resolution or our LC method, and the sodiated adduct ion provided better 472 quantification on the samples. We observed that the absolute differences in mass and CCS were 473 larger for IPP/DMAPP and GPP ions compared to the rest of the identified molecules 474 (respectively, -0.0858 and -0.0449 m/z, and 1.03 and 0.9 CCS). However, after adjusting these 475 values we could quantify these molecules across all samples and observed a consistent trend. 476 The levels of IPP/DMAPP, GPP (precursor to FPP) and extracellular FPP-derived bisabolene 477 (Fig. 7) which were higher in cells grown on HALM hydrolysates at 60 hr compared to all other 478 conditions, could explain the decreased level of HMGR detected in the proteomic analysis and 479 the subsequent accumulation of HMG-CoA (i.e., at 60 hr for the comparison of HALM vs HAHM, 480 log2FC of HMGR: -1.16, log2FC of HMG-CoA: 4.67, respective p-values are 1.82x10⁻³ and 481 3.28x10⁻³; see Supplementary Table 5). Previously, it had been observed that when sterol 482 pathway flux is high, degradation of HMGR is fast and its levels are low⁴⁶ and this is what was 483 revealed for GB 2 growth on HALM hydrolysate after 60 hours by our advanced analytical 484 workflow.

485 **Discussion**

486 Using synthetic biology applications, we have described and demonstrated an optimized 487 analytical method, a chromatography method prediction tool, and an alternative metabolomics 488 algorithm for robust processing of multidimensional data acquired in state-of-the-art LC-IM-MS 489 instrumentation. The advantages of using LC-IM-MS with DIA and PeakDecoder enable high-490 throughput analyses with increased metabolite coverage and more confident annotation due to 491 several aspects: 1) in terms of data acquisition, our 9 min LC method is faster than the GC 492 methods typically used, 2) the IM dimension further separates more analytes and increases 493 annotation confidence by combining CCS and RT compared to LC alone, 3) DIA further increases 494 annotation confidence with fragmentation information and provides better reproducibility and 495 dynamic range than DDA, and 4) Our PeakDecoder score provides a confident metric for 496 metabolite annotation. These tools have the potential to enable faster and more accurate testing 497 of strains generated by high-throughput engineering workflows and therefore accelerate the DBTL

498 cycle. Engineered microbes (e.g., bacteria, yeast, fungi) producing bioproducts (e.g., fuels, 499 chemicals, materials) in a sustainable way are necessary to achieve a strong bioeconomy and 500 decrease dependence on fossil fuels. Our analytical and computational workflow will provide 501 capabilities for fast analysis of current and new metabolites of interest and is broadly useful, 502 beyond the ABF consortium, in a wide range of environmental and biological metabolomics 503 research.

504

505 Our multidimensional metabolite library built from 64 standards is available as a resource to the 506 community and we expect it to be expanded, since the combination of RT and CCS with co-elution 507 and co-mobility profiles from DIA fragmentation patterns significantly increases confidence in 508 overall compound identification. Besides, DIA spectra are a permanent and comprehensive digital 509 record of all detectable ions in the sample, which can be re-processed as new libraries or new 510 tools become available, and without the need of reanalyzing the samples for acquiring new data. 511 Consequently, new bio-chemical hypothesis could be investigated using the existing microbial 512 data, and DIA would allow evaluating/investigating side effects such as undesired pathways 513 activated or undesired products, which would be missed with targeted-MS methods.

514

515 Although more developments could be explored, for example, other decoy generation methods, 516 training models specific to the number of fragments, engineering of ML features, comparing other 517 ML methods and evaluating other MS/MS similarity metrics such as the spectral entropy⁶, our 518 present results show better performance over existing LC-IM-MS tools for confident metabolite 519 annotation with PeakDecoder using ML features based on summary statistics and an SVM 520 classifier. Limitations of the current algorithm include requirements for sufficient high-guality peak-521 groups for training (i.e., limited performance for samples with very low complexity) and a library 522 acquired with compatible analytical conditions for inference.

523

524 Since the training strategy in PeakDecoder is to learn how to distinguish good co-elution and co-525 mobility patterns from the raw data directly and does not rely on fragmentation rules, its application 526 is not limited to a particular omics. We believe that PeakDecoder represents a step towards 527 universal software for molecular identification and it will potentially enable error rate calculations 528 for different analyte types. Future work will be performed to compare PeakDecoder to DDA 529 analyses and to evaluate it with predicted MS/MS, CCS and RT metabolomics libraries, as well 530 as applications for proteomics and lipidomics. While PeakDecoder was built on several MS-tools, 531 we envision a fully automated pipeline which is enhanced by replacing with novel artificial 532 intelligence (AI)-based methods the traditional tools that heavily require intervention from human 533 experts. Similarly to other research fields, advanced AI MS-tools may achieve human-level or 534 super-human AI systems⁴⁷ and have the potential to exploit the rich multidimensional LC-IM-MS 535 data to derive new molecular knowledge.

536

537 Methods

538 Automated chromatographic method selection software

539 The Automated chromatographic Method Selection Software (AMSS) used PubChem IDs as input 540 for information on molecules such as SMILES and physical and chemical properties, and utilized 541 the previously published BioCompound Machine Learning (BCML) tool⁴⁸ to calculate additional 542 physico-chemical properties. PaDEL descriptor⁴⁹ was used to compute molecular descriptors 543 which are used as features for further ML applications. As the training data provided for 544 developing the ML application was limited, the feature selection method Boruta⁵⁰ was applied to 545 avoid overfitting the predictive model. The random forest method was applied in sci-kit-learn for 546 ML predictive model following the feature selection. Datasets from previous HPLC analysis of 547 different compounds from the IROA Compound Test Set (SIGMA Chemical) were analyzed in ESI 548 positive and negative modes using four different chromatography methods (HILIC+, HILIC-, PR+, 549 RP-) and used as training: 467 compounds for pH 9.2 and 508 compounds for pH 2.7. The training 550 datasets had scores for performances of all analysis methods for different compounds. Using the 551 strategy mentioned above, four different categorical predictive models were built for each 552 chromatographic analysis method. The predictive model is used to predict the best 553 chromatograph analysis method for testing compounds. Additionally, Local Interpretable Model-554 Agnostic Explanations (LIME)⁵¹ was applied for model application explanation. LIME scores were 555 also used to draw shapes with color codes to highlight the chemical structural/substructure of the 556 compounds with prediction. This software is used to run all four predictive models on the new test 557 case. The predictions for all the four different models are reported as predictions. It should be 558 noted that there can be multiple suitable methods for HPLC analysis for a single compound.

559 Sample preparation

560 Standards: 64 commercially available compounds from the central carbon metabolism (common 561 to all ABF hosts) or metabolites that are part of pathways that had been engineered in the ABF 562 mutant strains were selected. Standards were prepared individually at a concentration of 25 μ M 563 using 3:2 acetonitrile: water as solvent. Once analyzed individually, standard mixes containing 564 10-15 metabolites at the same final concentration and solvent composition were prepared and 565 acquired in the analytical platforms.

566

567 A. pseudoterreus and A. niger strains: The A. pseudoterreus codon optimized β -alanine pathway 568 was detailed in our previous manuscript²⁴. The β -alanine pathway was randomly integrated into 569 A. niger ATCC 11414. Three transgenic strains (3HP-10, 5, and 9) producing low, medium, and 570 high levels of 3HP were selected for metabolite profiling. In addition, two transgene 571 overexpression constructs were built. A. niger aspartate aminotransferase (aat, Genebank 572 access: EHA22111.1) cDNA was under the control of A. niger translation elongation factor-1a 573 (tef1) promoter and its first intron and the transcriptional terminator of A. niger phosphoglycerate 574 kinase (pgk1), while A. niger pyruvate carboxylase (pyc, Genebank access: AJ009972.1) cDNA 575 was under the control of A. niger multiprotein-bridging factor-1 promoter and the transcriptional 576 terminator of pgk1. Both of transgene expression constructs were separately introduced into strain

577 3HP-9 to generate a series of new transgenic strains: 3HP-9 *aat*-1 to 12 or 3HP-9 *pvc*-1 to 12. 578 Transgenic strains 3HP-9 aat-5 and 3HP-9 pyc-1 producing higher and highest levels of 3HP 579 were selected for metabolite profiling. The selected strains were grown in 50 ml of the modified 580 Riscaldati B medium²⁴ in 250 ml PYREX Erlenmeyer flasks. The flasks were incubated at 30 °C 581 while shaking at 200 rpm. The supernatants and biomass were collected at day 4. For each 582 culture, 2 ml of supernatant was filtered through a 0.2 µm syringe filter and 1 ml of biomass was 583 collected via vacuum filtration through 2 layers of EMD Millipore miracloth and washed with 2 ml 584 of phosphate-buffer saline. The biomass was transferred into 1.5 ml microcentrifuge tubes and 585 immediately frozen in liquid nitrogen. Both supernatants and biomass pellets were stored at -80 586 °C prior to extraction of metabolites.

587

588 P. putida strains: Detailed explanation about the integration site selection, plasmid design, 589 assembly and transformation were presented previously³⁷. Briefly, P. putida KT2440 was used as 590 the wild-type strain. An mKate fluorescent reporter construct was designed, synthesized, and 591 introduced into seven different insertion locations on the P. putida KT2440 chromosome by 592 homologous recombination, always in the same orientation. Growth and fluorescence of these 593 seven mKate expression strains were measured in M9 minimal medium containing 30 mM 594 glucose and reported in the cited manuscript. To further test the effect of the integration locus on 595 function of a heterologous gene, a functional protein (muconate transporter) was integrated into 596 three of the seven sites and the resulting variation in growth and protein expression was 597 measured. The selection of the integration sites chosen for additional characterization was based 598 on their display of different phenotypes with mKate, such as slow growth and low fluorescence 599 (PP 2224), slow growth and high fluorescence (PP 1642), or WT growth and medium fluorescence 600 (PP 5042). Overnight cultures of WT P. putida KT2440 and strains carrying a codon-optimized 601 copy of mucK (the gene coding for the muconate importer) were inoculated into 10 mL LB medium 602 to give a starting culture density of 0.2 OD600 nm and were incubated at 30°C with shaking until 603 the culture density reached 1.0 OD600 nm. Cell cultures were centrifuged and washed twice in 604 M9 salts before resuspending in the same buffer. The washed cells were used to inoculate 50 mL 605 M9 medium containing 30 mM glucose (for WT KT2440) or 30 mM cis, cis-muconate (for strains 606 containing mucK insertions). The starting culture density was 0.1 OD600 nm and growth 607 continued until 0.7 OD600 nm was reached. Cells were collected by centrifugation and were 608 washed one time with ice-cold PBS. Cell pellets were weighed (~50 mg of wet weight collected), 609 flash frozen in liquid nitrogen and stored at -80 °C prior to shipment and extraction of metabolites 610 and proteins. Omics samples were prepared in triplicate.

611

612 R. toruloides strains: The R. toruloides strain used in this study, GB2, was described in detail in 613 our previous manuscript⁴². Its parent strain, BIS3, was the highest bisabolene producer of a panel 614 of P_{GAPDH} -BIS strains that were modified only by insertion of a heterologous α -bisabolene synthase gene (BIS) from Abies grandies under control of the native R. toruloides GAPDH 615 616 (glyceraldehyde 3-phosphate dehydrogenase) promoter into WT R. toruloides and differed in copy 617 number only^{40,42}. BIS3, was selected for addition of a second expression cassette consisting of 618 BIS under control of the native R. toruloides ANT (adenine nucleotide translocase) promoter, 619 which resulted in strain GB2. GB2 contains 6 copies of the PANT-BIS cassette in addition to the 620 original 10 copies of the P_{GAPDH}- BIS cassette in BIS3. GB2 cells were grown in vessels in an 621 Ambr® 250 High Throughput system (Sartorius) with a total volume of 150 ml each. The growth 622 media consisted of four DMR (deacetylation and mechanical refining method) hydrolysates made 623 from corn stover by the National Renewable Energy Laboratory (Golden, Colorado) from a 2x2 624 matrix of ash (high/low) and moisture (high/low). The DMR hydrolysates were referred to as HALM 625 (high ash, low moisture), HAHM (high ash, high moisture), LALM (low ash, low moisture) and 626 LAHM (low ash, high moisture). The media were only supplemented with ammonium sulfate (5.00 627 q/L), potassium phosphate monobasic (10.34 q/L) and potassium phosphate dibasic (4.18 q/L), 628 pH was controlled at 5.0 by addition of ammonium hydroxide. Dissolved oxygen was set as 30%, 629 air flow 75 standard liter per minute (= 0.5 volume of air sparged in aerobic cultures per unit 630 volume of growth medium per minute), agitation (cascade) of 500-2000 rpm and growth 631 temperature of 30°C. A dodecane overlay (20% of total volume) was added to capture the 632 bisabolene produced. Three bioreactors were prepared for each condition (hydrolysate). For 633 omics measurements, five mL volume of culture were taken from each Ambr fermentation vessel 634 at time points 24 and 60 hrs and centrifuged at 4,000xg at 4°C for 5 min. The supernatant and 635 dodecane overlay were decanted and transferred to another tube for bisabolene analysis, done 636 by GC-MS as described previously⁵². The cell pellet was resuspended in 1.5 ml of ice-cold PBS 637 and transferred to a new tube. Samples were centrifuged for 5 min at 16,000xg, the PBS removed, 638 and the cell pellet was flash frozen with liquid nitrogen. Pellets were stored at -80C until shipment 639 and extraction of metabolites and proteins.

640

641 All microbial samples (cell pellets) were extracted using the MPLEx protocol as previously 642 reported^{24,37,53}. Briefly, a mixture of chloroform, methanol and water was added to the cell pellets, 643 extraction done in an ice bath and the polar and non-polar phases were combined and dried under 644 vacuum. Dried extracts were resuspended in 300 μ l of 3:2 acetonitrile:water, transferred to an 645 LC-MS vial and stored at -20°C until analysis.

646 SRM and LC-IM-MS analyses

647 Ultrahigh performance liquid chromatography (UHPLC) methods were implemented and 648 optimized by analyzing standards. Chromatographic separation was performed with an Agilent 649 UHPLC 1290 Infinity II system. The sample injection volume was 10 µL and the autosampler 650 temperature was maintained at 4°C. The Agilent UHPLC was equipped with a Water XBridge 651 BEH Amide XP Column, 2.5 µm (2.1 mm i.d. X 50 mm). A Waters XBridge BEH Amide XP 652 VanGuard cartridge, 2.5 µm (2.1 mm i.d. X 5 mm) was also installed to remove potential 653 particulate contamination from the mobile phases. Mobile phases consisted of (A) 10 mM 654 ammonium acetate, 10 µM InfinityLab deactivator additive, pH 9.2 in 90% water and 10 % 655 acetonitrile, (B) 10 mM ammonium acetate, pH 9.2 in 90% acetonitrile. The column was kept at 50°C throughout the run. The gradient length was 8.70 min (detailed as following, 0.0:0.350:90. 656 657 1.0:0.350:90, 1.1:1.0:85, 4.0:0.750:80, 5.0:0.750:40, 6.5:0.750:40, 6.8:0.750:20, 7.0:0.750:20, 658 7.5:0.750:90 in terms of min:flow-rate-µL/min:%B) with an equilibration time of 3.0 min. The 659 UHPLC system was coupled to an Agilent 6490 triple quadrupole (QQQ) for initial method 660 development. Scan and SRM analyses were performed for precursor fragmentation and transition 661 identification. The instrument was operated in the negative polarity with the following parameters: 662 ion spray voltage of 3000 V, capillary inlet temperature of 225°C, gas flow 15 ml/min, nebulizer

663 pressure 20 psi, sheath gas temperature 250°C, sheath gas flow 11 ml/min. Data were acquired 664 in a mass range from 65 to 1400 m/z. SRM analyses were also performed for the calibration 665 curves of example standards and the evaluation in microbial samples. Data was processed in 666 Skyline¹⁴ (v.64.21.1.0.146) for peak area integration. A total of 11 samples were analyzed by SRM 667 for the *P. putida* strains, with 3 biological replicates for all, except mucK PP2224 that had 2).

668

669 The optimized UHPLC system was coupled to an Agilent 6560 Drift Tube Ion Mobility 670 Spectrometry (DTIMS)-QTOF MS (Agilent Technologies, Santa Clara, CA). The MassHunter data 671 acquisition software (v.B.09.00 (B9044.0), Agilent Technologies) was used to collect all mass 672 spectrometry raw data files. The instrument was mass-calibrated before every batch 673 measurement using the Agilent ESI Tune solution. Standard mixes and microbial samples were 674 analyzed in negative mode using a Dual AJS ESI and high-purity nitrogen as drift gas. Parameters 675 were set to 325 degrees C gas temperature, 5L/min drying gas, 30psi nebulizer, 275 degrees C 676 sheath gas, 11L/min sheath gas flow, 2500V Vcap, 2000V nozzle voltage and 400V fragmentor. 677 Data was acquired in All-lons DIA mode alternating between low (MS1) and high (MS2) collision 678 energies at the frequency of 2 frames per second. 60 ms of maximum drift time was allowed with 679 19 transients per frame. Mass range 50-110 m/z. Fixed CE values of 20 or 40 V were used to 680 cover both labile and compounds with masses > 600 Da. A total of 81 microbial samples were 681 analyzed by LC-IM-MS: 46 for the A. pseudoterreus and A. niger strains (4 biological replicates 682 for each condition, except groups Control (Exp 1, A. pseudoterreus cad) and F (Exp. 1. A. niger 683 3-HP high) that had 3 each; analyzed with 20 V CE only), 11 for the P. putida strains (3 biological 684 replicates for all, except muck PP2224 that had 2; analyzed with 20 and 40 V CEs), and 24 for 685 the *R. toruloides* strains (4 biological replicates for each group; analyzed with 20 and 40 V CEs).

686 Data processing for LC-IM-MS

687 CCS were calculated using the IM-MS Browser (v.10.0, Agilent Technologies) with the single-field 688 method⁵⁴ and the Agilent Tune-Mix solution as calibrants. The PNNL-Preprocessor⁵⁵ 689 (v2020.07.24) was used to apply moving average smoothing (points: 3 in LC and 3 in IM) and 690 filtering (minimum intensity threshold 20 counts). MS-DIAL¹¹ (v.4.70) was used to perform 691 untargeted feature finding and MS/MS deconvolution (soft ionization, ion mobility separation, data 692 independent MS1, MS and MS/MS profile data, negative ion mode, metabolomics, centroid MS1 693 tolerance 0.01, centroid MS2 tolerance 0.025, smoothing level 1, minimum peak width 3, minimum 694 peak height 300, peak spotting mass slice width 0.1, deconvolution sigma window 0.5, MS2Dec 695 amplitude cut off 0, alignment RT tolerance 0.1, alignment MS1 tolerance, 0.015, alignment RT 696 factor 0.5 and ion mobility accumulated RT range 0.2). Skyline¹⁴ (v.64.21.1.0.146) was used to 697 perform targeted data extraction (acquisition method DIA, isolation scheme All lons, mass 698 analyzer TOF, mass resolving power 10000, ion mobility resolving power 40 and small molecule 699 fragment types "p,f"). Implementation of the PeakDecoder algorithm and evaluation of the results 700 were performed in R (v.4.1.0, A language and environment for statistical computing, R Foundation 701 for Statistical Computing, Vienna, Austria, https://www.R-project.org), using packages e1071 702 (v.1.7-9) and ggplot2 (v.3.3.3).

703

704 The following approximation was used to calculate the negative mobility offset of the fragments 705 from their precursors: ((FragmentMz - PrecursorMz)/ PrecursorMz)*0.7 - PrecursorMz*0.0001. 706 Which overall worked well for both 20V and 40V collision energies and resulted in values mostly 707 between -0.1 and -0.3 msec, with smaller m/z ions showing larger offsets. This negative drift time 708 shift is a function of the collision energy used and the mass of the fragment ion and it can be 709 explained by the fact that under the accelerating electric field smaller ion fragments move faster 710 through the collision cell and the ion beam compressor region during high-energy steps than 711 larger precursor ions³³; hence t0, i.e. the time ions spend traveling though the instrument, outside 712 the drift tube, is different.

713

A library with precursor *m/z*, RT, CCS and fragment *m/z* values was built from the standards. RT and transitions were obtained from the SRM results. CCS and additional transitions were determined post-acquisition from the in the LC-IM-MS data. The list of 64 metabolites with accurate mass, RT and CCS are in Supplementary Table 1. The list of fragments in csv format and the full library in the NIST MSP text format library for metabolite identifications are in Supplementary Data 1.

720 Metabolite scoring for LC-IM-MS: PeakDecoder algorithm

<u>1) Feature finding and fragment ion deconvolution:</u> data is processed in untargeted mode (MS-DIAL) to extract all precursor ion features (MS1) and their respective deconvoluted fragment ions (MS2) based on co-elution and co-mobility. The alignment (Peak ID matrix, msp format) and all peak lists (txt, centroid) are exported.

725

<u>2) Target and decoy generation:</u> an R script was implemented to generate a training set. The MS DIAL alignment results including features and their fragments is used as input. Features with S/N
 >= 15 and at least 3 fragments with intensities within 1-130% of their precursor intensity are kept
 as targets. The top 16 most intense fragments are kept per target.

- To generate the decoys, the set of targets are associated by pairs. For each target, another target is found from the same representative LC-IM-MS run, which the precursor m/z is within 50 units difference (to ensure that the paired features are from molecules of similar size), has the largest RT difference (at least 3 min to avoid pairing a repeated feature from a large tailing peak) and has the same number of fragments. A pair of decoys is generated for the paired targets by keeping the same precursor properties and swapping the m/z values of 40-60% of the fragments randomly chosen from the top-most-intense. Targets for which decoys could not be generated are excluded.
- A transition list in Skyline format is generated with this preliminary set of targets and decoys.
- 738
- <u>3) Targeted data extraction:</u> the transition list for the training set and the query metabolites in the
 library are processed separately. The precursor and fragment ion signals are extracted (Skyline)
 from all the LC-IM-MS runs. The two reports (training set and query metabolites) are exported,
 which include the required XIC metrics: area, height, mass error, FWHM (LC), RT, expected RT,
- 743 expected CCS.
- 744

745 4) Machine learning training: an R script was implemented for training. The Skyline report of the 746 preliminary training set is used as input. The targets are filtered according to multiple thresholds 747 to ensure a good quality training set. Fragments with unassigned height (i.e., NA or zero) and 748 precursors with S/N < 20 are removed. Each fragment is evaluated to count the number of low-749 quality metrics: area <= 0, height < 1% their precursor height, mass error > 15 ppm, RT difference 750 to their precursor larger than 0.1 min and FWHM difference to their precursor larger than 2x the 751 precursor FWHM. Targets with at least 2 fragments with high-quality metrics are kept. To simulate 752 interferences some fragments with low-quality metrics are kept. Fragments with the worst metrics 753 and ranked higher than 2x the number of fragments with good metrics are removed. For each LC-754 IM-MS run, only the paired decoys with the same subset of fragments as their targets after filtering 755 are kept for maintaining the same distribution of targets and decoys by number of fragments and 756 m/z values. The target fragment height is used as the expected intensity and assigned to the 757 corresponding decoys to minimize the impact of peak integration differences between MS-DIAL 758 and Skyline. The filtered targets with at least 3 fragments in total and their corresponding decoys 759 are used as the final training set. The following descriptors are calculated for the filtered training 760 set and used as ML features:

- 761 762
- DIA-cosSim: cosine similarity between the integrated area and the expected intensity of the fragments.
- DIA-RTdiffSd and DIA-RTdiffMean: standard deviation and mean of the differences
 between the precursor RT and RT of its fragments.
- DIA-FWHMdiffSd and DIA-FWHMdiffMean: standard deviation and mean of the differences between the precursor FWHM and FWHM of its fragments.
- DIA-MassErrorSd and DIA-MassErrorMean: standard deviation and mean of the fragment mass errors.
- 769

770 An SVM binary classifier (e1071 R package) is trained using a radial kernel, scaling (to zero mean 771 and unit variance), 10-fold cross validation and probability calculation. The probability is 772 calculated by fitting a logistic distribution using maximum likelihood to the decision values of all 773 binary classifiers, and computing the a-posteriori class probabilities for the multi-class problem 774 using quadratic optimization. The trained model is saved. The target probabilities are calculated 775 for the full training set and a confusion matrix and FDR are calculated to evaluate performance. The FDR is calculated as FP/(TP + FP)⁵⁶. The target probability is used as the PeakDecoder 776 777 score. A table with pairs of values (FDR, PeakDecoder score) is automatically generated after 778 training (file PeakDecoder-FDR-thresholds [dataset].csv).

779

780 5) Machine learning inference: an R script was implemented for inference. The model previously 781 trained and saved is loaded. The Skyline report for the query metabolites and the library with the 782 fragment ion abundances generated from the standards are used as input. The descriptors are 783 calculated as described above. The precursor RT error (minutes) is calculated as the difference 784 between the run RT and the expected RT (from the standards). The CCS error is calculated as 785 the percentage difference between the run CCS (obtained from the corresponding MS-DIAL peak 786 lists, since Skyline uses the CCS as a filter and does not report the actual CCS from the IM peak 787 apex in the run) and the expected CCS (from the standards). A query metabolite is considered identified if, in at least one of the runs, passes all cutoff thresholds: precursor mass error < 18 788

ppm, precursor RT error < 0.4 min, CCS error < 0.8 % and PeakDecoder score > 0.8 (or
corresponding to 1% FDR).

191

792 Pathway analyses

793 Metabolites with at least 1 replicate identified with high confidence were selected and their 794 Skyline-integrated precursor and fragment ion abundances across all runs were used for 795 guantitation. Statistical analysis of the metabolite abundance data was performed in R using the 796 pmartR package⁵⁷ (v0.9.0). For *P. putida* and *R. toruloides* datasets, the mean values of 797 abundances acquired with 20V and 40V CE were used for the analysis. The abundance values 798 were log2 transformed, and the test for differential abundance between control and test samples 799 was performed using the IMD-ANOVA method⁵⁸. Clustered heatmaps of log2 abundances were 800 generated using the R package pheatmap (v1.0.12) with Euclidean distance and complete linkage 801 method. Bar and error bar plots shown on the metabolic pathway maps were generated using the 802 python package matplotlib (v3.5.1). The metabolic pathway maps for A. pseudoterreus/A. niger 803 and *R. toruloides* were drawn based on the genome-scale metabolic models iJB1325⁵⁹ and 804 Rt IFO0880⁶⁰ using ChemDraw (v19.0). The metabolomics and proteomics data visualization on 805 the P. putida metabolic pathway map was performed with the genome-scale metabolic model 806 iJN1462⁶¹ using the python packages escher⁶² (v1.7.3) and cobrapy⁶³ (v0.22.1).

807 GC-MS analyses

808 Dried extracts for metabolomics analysis were obtained after MPLEx extraction as explained in 809 the Sample Preparation section. The stored metabolite extracts were completely dried under 810 speed-vacuum to remove moisture and chemically derivatized as previously reported⁶⁴. Briefly, 811 the extracted metabolites were derivatized by methoxyamination and trimethylsilyation (TMS), 812 then the samples were analyzed by GC-MS. Samples were run in an Agilent GC 7890A using a 813 HP-5MS column (30 m × 0.25 mm × 0.25 µm; Agilent Technologies, Santa Clara, CA) coupled 814 with a single guadrupole MSD 5975C (Agilent Technologies). One microliter of sample was 815 injected into a splitless port at constant temperature of 250°C. The GC temperature gradient 816 started at 60 °C and hold for 1 minute after injection, followed by increase to 325 °C at a rate of 817 10 °C/minute and a 5-minute hold at this temperature. Fatty acid methyl ester standard mix (C8-818 28) (Sigma-Aldrich) was analyzed in parallel as standard for retention time calibration. GC-MS 819 raw data files were processed using the Metabolite Detector (v2.5)⁶⁵. Retention indices (RI) of 820 detected metabolites were calculated based on the analysis of a FAMEs mixture, followed by their 821 chromatographic alignment across all analyses after deconvolution. Metabolites were initially 822 identified by matching experimental spectra to a PNNL augmented version of Agilent GC-MS 823 metabolomics Library, containing spectra and validated retention indices for over 850 metabolites. 824 Then, the unknown peaks were additionally matched with the NIST17/Wiley11 GC-MS library. All 825 metabolite identifications and quantification ions were validated and confirmed to reduce 826 deconvolution errors during automated data-processing and to eliminate false identifications. A 827 total of 46 samples of the A. pseudoterreus and A. niger strains were analyzed by GC-MS, with 4

biological replicates for each condition, except groups Control (Exp 1, A. pseudoterreus cad) and
F (Exp. 1. A. niger 3HP high) that had 3 each.

830 Targeted proteomics analyses of *P. putida* strains

831 Intracellular proteins from samples of *P. putida* strains, KT2440, with heterologous gene insertion 832 were extracted and digested as previously described³⁷. Peptides from previously established assavs³⁹ were used for the targeted proteomics analysis of enzymes in various metabolic 833 834 pathways. Analysis of the targeted proteomics assay was performed via LC-SRM. To facilitate 835 protein quantification, crude heavy peptide mixture stock solution was spiked in the 0.20 µg/µL 836 peptide samples at a nominal concentration of 25 fmol/µL for each peptide. LC-SRM analysis 837 utilized a nanoACQUITY UPLC® system (Waters Corporation, Milford, MA) coupled online to a 838 TSQ Altis triple quadrupole mass spectrometer (Thermo Fisher Scientific). The UPLC® system 839 was equipped with an ACQUITY UPLC BEH 1.7 µm C18 column (100 µm i.d. × 10 cm) and the 840 mobile phases were (A) 0.1% formic acid in water and (B) 0.1% formic acid in acetonitrile. 2 µL 841 of sample (i.e., 0.4 µg of peptides) were loaded onto the column and separated using a 110-min 842 gradient profile as follows (min:flow-rate-µL/min:%B): 0:0.4:1, 6:0.6:1, 7:0.4:1, 9:0.4:6, 40:0.4:13, 843 70:0.4:22, 80:0.4:40, 85:0.4:95, 91:0.5:95, 92:0.5:95, 93:0.5:50, 94:0.5:95, 95:0.6:1, 98:0.4:1. The 844 LC column was operated at a temperature of 42 °C. The TSQ Altis triple quadrupole mass 845 spectrometer was operated with ion spray voltages of 2100 ± 100 V and a capillary inlet 846 temperature of 350 °C. Tube lens voltages were obtained from automatic tuning and calibration 847 without further optimization. Both Q1 and Q3 were set at unit resolution of 0.7 FWHM and Q2 gas 848 pressure was optimized at 1.5 mTorr. The transitions were scanned with a dwell time of 0.78 849 msec. Targeted proteomics data were imported into Skyline (v64.22.2.1.278)⁶⁶ and the peak 850 boundaries were manually inspected to ensure correct peak assignment and peak boundaries. 851 Peak detection and integration were determined based on two criteria: 1) the same LC retention 852 time and 2) approximately the same relative peak intensity ratios across multiple transitions 853 between the light peptides and heavy peptide standards. The total peak area ratios of 854 endogenous light peptides and their corresponding heavy isotope-labeled internal standards from 855 Skyline were used for subsequent protein abundance rollup and pathway analysis.

856 Global proteomics analyses of *R. toruloides* strains

857 Intracellular proteins from samples of bisabolene producing R. toruloides strains, GB2.0, were 858 extracted, digested with trypsin, and analyzed by LC-MS/MS following a previously established 859 protocol⁶⁰. A Q-Exactive Plus Orbitrap mass spectrometer (Thermo Fisher Scientific) was used in this study with the parameters as following: full MS (AGC, 3×10⁶; resolution, 70,000; m/z range, 860 861 300–1800; maximum ion time, 20 ms); MS/MS (AGC, 1×10⁵; resolution, 17,500; m/z range, 200– 862 2000; maximum ion time, 50 ms; minimum signal threshold, 5×10³; isolation width, 1.5 Da; 863 dynamic exclusion time setting, 30 s; collision energy, NCE 30; TopN, 12). The MS data were 864 searched against the R. toruloides strain IFO0880 (v4.0) and heterologous protein sequences [https://mycocosm.igi.doe.gov/Rhoto IFO0880 4]⁴⁵ using MaxQuant⁶⁷ (v1.6.2.10) and the 865 866 following parameters: 1% peptide-level and protein-level FDR, match-between-runs enabled, 867 partial tryptic with trypsin/P, maximum missed cleavage of 2, dynamic modification of oxidation

on methionine and N-terminal acetylation, fixed carbamidomethyl on cysteine, mass tolerances
of 20 ppm for both precursor and fragment ions, minimum peptide length of 7, and a minimum
number of unique peptides for protein quantification as 1. Peptide intensity level data from
MaxQuant were further processed by pmartR (v0.9.0) for quality control, protein rollup, and
statistical comparisons.

873 Data availability

874 The microbial LC-IM-MS data (and related Skyline projects) generated in this study have been 875 deposited in the MassIVE database under accession code MSV000089733 876 [https://doi.org/doi:10.25345/C52R3P17Z]. The P. putida targeted proteomics data generated in 877 this study have been deposited in the Panorama database [https://doi.org/10.6069/6j7y-t592]. 878 The R. toruloides global proteomics data generated in this study have been deposited in the 879 MassIVE database under accession code MSV000091202 880 [https://doi.org/doi:10.25345/C50K26N04]. Source data are provided with this paper.

881 Code availability

882 The source code of PeakDecoder⁶⁸, the library built from standards, and all the input files and 883 results can be found at https://github.com/EMSL-Computing/PeakDecoder. The source code of 884 the automated chromatographic method selection software can be found at 885 https://github.com/poorey/AMSS.

886 **References**

- Liebal, U. W., Phan, A. N. T., Sudhakar, M., Raman, K. & Blank, L. M. Machine Learning
 Applications for Mass Spectrometry-Based Metabolomics. *Metabolites* 10,
 doi:10.3390/metabo10060243 (2020).
- 890 2 Gowda, G. A. & Djukovic, D. Overview of mass spectrometry-based metabolomics:
 891 opportunities and challenges. *Methods Mol Biol* **1198**, 3-12, doi:10.1007/978-1-4939892 1258-2 1 (2014).
- 893 3 Hillson, N. *et al.* Building a global alliance of biofoundries. *Nat Commun* **10**, 2040, doi:10.1038/s41467-019-10079-2 (2019).
- Chaleckis, R., Meister, I., Zhang, P. & Wheelock, C. E. Challenges, progress and
 promises of metabolite annotation for LC-MS-based metabolomics. *Curr Opin Biotechnol*55, 44-50, doi:10.1016/j.copbio.2018.07.010 (2019).
- Shang, X. W., Li, Q. H., Xu, Z. D. & Dou, J. J. Mass spectrometry-based metabolomics
 in health and medical science: a systematic review. *Rsc Adv* 10, 3092-3104,
 doi:10.1039/c9ra08985c (2020).
- Building State
 <
- Burnum-Johnson, K. E. *et al.* Ion Mobility Spectrometry and the Omics: Distinguishing
 Isomers, Molecular Classes and Contaminant Ions in Complex Samples. *Trends Analyt Chem* 116, 292-299, doi:10.1016/j.trac.2019.04.022 (2019).

907 8 Bilbao, A. et al. Processing strategies and software solutions for data-independent 908 acquisition in mass spectrometry. Proteomics 15, 964-980, doi:10.1002/pmic.201400323 909 (2015). 910 9 Gillet, L. C. et al. Targeted data extraction of the MS/MS spectra generated by data-911 independent acquisition; a new concept for consistent and accurate proteome analysis. 912 Mol Cell Proteomics 11, O111 016717, doi:10.1074/mcp.O111.016717 (2012). 913 10 Guo, J. & Huan, T. Comparison of Full-Scan, Data-Dependent, and Data-Independent 914 Acquisition Modes in Liquid Chromatography-Mass Spectrometry Based Untargeted 915 Metabolomics. Anal Chem 92, 8072-8080, doi:10.1021/acs.analchem.9b05135 (2020). 916 Tsugawa, H. et al. A lipidome atlas in MS-DIAL 4. Nat Biotechnol 38, 1159-1163, 11 917 doi:10.1038/s41587-020-0531-2 (2020). 918 12 Chen, G. et al. Customized Consensus Spectral Library Building for Untargeted 919 Quantitative Metabolomics Analysis with Data Independent Acquisition Mass 920 Spectrometry and MetaboDIA Workflow. Anal Chem 89, 4897-4906. 921 doi:10.1021/acs.analchem.6b05006 (2017). 922 13 Guo, J., Shen, S., Xing, S. & Huan, T. DaDIA: Hybridizing Data-Dependent and Data-923 Independent Acquisition Modes for Generating High-Quality Metabolomic Data. Anal 924 Chem 93, 2669-2677, doi:10.1021/acs.analchem.0c05022 (2021). 925 14 MacLean, B. X. et al. Using Skyline to Analyze Data-Containing Liquid Chromatography, 926 Ion Mobility Spectrometry, and Mass Spectrometry Dimensions. J Am Soc Mass 927 Spectrom 29, 2182-2188, doi:10.1007/s13361-018-2028-5 (2018). 928 Li, H., Cai, Y., Guo, Y., Chen, F. & Zhu, Z. J. MetDIA: Targeted Metabolite Extraction of 15 929 Multiplexed MS/MS Spectra Generated by Data-Independent Acquisition. Anal Chem 88, 930 8757-8764, doi:10.1021/acs.analchem.6b02122 (2016). 931 Alka, O. et al. DIAMetAlyzer allows automated false-discovery rate-controlled analysis 16 932 for data-independent acquisition in metabolomics. Nat Commun 13, 1347, 933 doi:10.1038/s41467-022-29006-z (2022). 934 17 Stancliffe, E., Schwaiger-Haber, M., Sindelar, M. & Patti, G. J. DecoID improves 935 identification rates in metabolomics through database-assisted MS/MS deconvolution. 936 Nat Methods 18, 779-787, doi:10.1038/s41592-021-01195-3 (2021). 937 18 Blazenovic, I. et al. Structure Annotation of All Mass Spectra in Untargeted Metabolomics. Anal Chem 91, 2155-2162, doi:10.1021/acs.analchem.8b04698 (2019). 938 939 19 Palmer, A. et al. FDR-controlled metabolite annotation for high-resolution imaging mass 940 spectrometry. Nat Methods 14, 57-60, doi:10.1038/nmeth.4072 (2017). 941 20 Scheubert, K. et al. Significance estimation for large scale metabolomics annotations by 942 spectral matching. Nat Commun 8, 1494, doi:10.1038/s41467-017-01318-5 (2017). 943 21 Wang, X. et al. Target-Decoy-Based False Discovery Rate Estimation for Large-Scale 944 Metabolite Identification. J Proteome Res 17, 2328-2334, 945 doi:10.1021/acs.jproteome.8b00019 (2018). 946 22 Li, D. et al. XY-Meta: A High-Efficiency Search Engine for Large-Scale Metabolome 947 Annotation with Accurate FDR Estimation. Anal Chem 92, 5701-5707, 948 doi:10.1021/acs.analchem.9b03355 (2020). 949 23 Kampers, L. F. C. et al. In silico-guided engineering of Pseudomonas putida towards 950 growth under micro-oxic conditions. Microb Cell Fact 18, 179, doi:10.1186/s12934-019-951 1227-5 (2019). 952 24 Pomraning, K. R. et al. Integration of Proteomics and Metabolomics Into the Design, 953 Build, Test, Learn Cycle to Improve 3-Hydroxypropionic Acid Production in Aspergillus 954 pseudoterreus. Front Bioeng Biotech 9, doi:ARTN 603832 955 10.3389/fbioe.2021.603832 (2021).

- 25 Lyu, L. *et al.* Engineering the Oleaginous Yeast Rhodosporidium toruloides for Improved
 p57 Resistance Against Inhibitors in Biomass Hydrolysates. *Front Bioeng Biotechnol* 9,
 p58 768934, doi:10.3389/fbioe.2021.768934 (2021).
- Berlanga-Clavero, M. V. *et al.* Bacillus subtilis biofilm matrix components target seed oil
 bodies to promote growth and anti-fungal resistance in melon. *Nat Microbiol* 7, 10011015, doi:10.1038/s41564-022-01134-8 (2022).
- 962 27 Bilbao, A. *et al.* Ranking Fragment Ions Based on Outlier Detection for Improved Label963 Free Quantification in Data-Independent Acquisition LC-MS/MS. *J Proteome Res* 14,
 964 4581-4593, doi:10.1021/acs.jproteome.5b00394 (2015).
- 865 28 Rosenberger, G. *et al.* Statistical control of peptide and protein error rates in large-scale
 866 targeted data-independent acquisition analyses. *Nat Methods* 14, 921-927,
 967 doi:10.1038/nmeth.4398 (2017).
- 968 29 Reiter, L. *et al.* mProphet: automated data processing and statistical validation for large-969 scale SRM experiments. *Nat Methods* **8**, 430-435, doi:10.1038/nmeth.1584 (2011).
- 97030Rost, H. L. et al. OpenSWATH enables automated, targeted analysis of data-971independent acquisition MS data. Nat Biotechnol 32, 219-223, doi:10.1038/nbt.2841972(2014).
- 31 Åhrne, E. *et al.* An improved method for the construction of decoy peptide MS/MS
 374 spectra suitable for the accurate estimation of false discovery rates. *Proteomics* 11, 4085-4095, doi:10.1002/pmic.201000665 (2011).
- 976 32 Cheng, C. Y., Tsai, C. F., Chen, Y. J., Sung, T. Y. & Hsu, W. L. Spectrum-based method
 977 to generate good decoy libraries for spectral library searching in peptide identifications. J
 978 Proteome Res 12, 2305-2310, doi:10.1021/pr301039b (2013).
- Mairinger, T. *et al.* Rapid screening methods for yeast sub-metabolome analysis with a
 high-resolution ion mobility quadrupole time-of-flight mass spectrometer. *Rapid Commun Mass Spectrom* **33 Suppl 2**, 66-74, doi:10.1002/rcm.8420 (2019).
- Blazenovic, I., Kind, T., Ji, J. & Fiehn, O. Software Tools and Approaches for Compound
 Identification of LC-MS/MS Data in Metabolomics. *Metabolites* 8,
 doi:10.3390/metabo8020031 (2018).
- Borodina, I. *et al.* Establishing a synthetic pathway for high-level production of 3hydroxypropionic acid in Saccharomyces cerevisiae via beta-alanine. *Metab Eng* 27, 5764, doi:10.1016/j.ymben.2014.10.003 (2015).
- 98836Nikel, P. I., Chavarria, M., Danchin, A. & de Lorenzo, V. From dirt to industrial989applications: Pseudomonas putida as a Synthetic Biology chassis for hosting harsh990biochemical reactions. Curr Opin Chem Biol 34, 20-29, doi:10.1016/j.cbpa.2016.05.011991(2016).
- 37 Chavés, J. E. *et al.* Evaluation of chromosomal insertion loci in the Pseudomonas putida
 37 KT2440 genome for predictable biosystems design. *Metab Eng Commun* **11**, e00139,
 394 doi:10.1016/j.mec.2020.e00139 (2020).
- 88 Kukurugya, M. A. *et al.* Multi-omics analysis unravels a segregated metabolic flux
 896 network that tunes co-utilization of sugar and aromatic carbons in Pseudomonas putida.
 897 J Biol Chem 294, 8464-8479, doi:10.1074/jbc.RA119.007885 (2019).
- 99839Gao, Y. et al. High-Throughput Large-Scale Targeted Proteomics Assays for Quantifying999Pathway Proteins in Pseudomonas putida KT2440. Front Bioeng Biotechnol 8, 603488,1000doi:10.3389/fbioe.2020.603488 (2020).
- Yaegashi, J. *et al.* Rhodosporidium toruloides: a new platform organism for conversion
 of lignocellulose into terpene biofuels and bioproducts. *Biotechnol Biofuels* 10, doi:ARTN
 241

1004 10.1186/s13068-017-0927-5 (2017).

100541Peralta-Yahya, P. P. *et al.* Identification and microbial production of a terpene-based1006advanced biofuel. Nat Commun 2, doi:ARTN 483

1007 10.1038/ncomms1494 (2011). 1008 Kirby, J. et al. Further engineering of R. toruloides for the production of terpenes from 42 1009 lignocellulosic biomass. Biotechnol Biofuels 14, doi:ARTN 101 1010 10.1186/s13068-021-01950-w (2021). 1011 Adams, P. in Greenhouse Gas Balances of Bioenergy Systems (eds Patricia Thornley 43 1012 & Paul Adams) 221-236 (Academic Press, 2018). 1013 44 Gardner, R. G. & Hampton, R. Y. A highly conserved signal controls degradation of 3-1014 hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) reductase in eukaryotes. J Biol Chem 274, 31671-31678, doi:DOI 10.1074/jbc.274.44.31671 (1999). 1015 1016 45 Coradetti, S. T. et al. Functional genomics of lipid metabolism in the oleaginous yeast 1017 Rhodosporidium toruloides. *Elife* 7, doi:ARTN e32110 1018 10.7554/eLife.32110 (2018). 1019 46 Garza, R. M., Tran, P. N. & Hampton, R. Y. Geranylgeranyl Pyrophosphate Is a Potent 1020 Regulator of HRD-dependent 3-Hvdroxy-3-methylglutaryl-CoA Reductase Degradation 1021 in Yeast. J Biol Chem 284, 35368-35380, doi:10.1074/jbc.M109.023994 (2009). 1022 47 Matsuo, Y. et al. Deep learning, reinforcement learning, and world models. Neural 1023 Networks, doi:https://doi.org/10.1016/j.neunet.2022.03.037 (2022). 1024 Whitmore, L. S. et al. BioCompoundML: A General Biofuel Property Screening Tool for 48 1025 Biological Molecules Using Random Forest Classifiers. *Energy & Fuels* **30**, 8410-8418, 1026 doi:10.1021/acs.energyfuels.6b01952 (2016). 1027 49 Yap, C. W. PaDEL-descriptor: an open source software to calculate molecular 1028 descriptors and fingerprints. J Comput Chem 32, 1466-1474, doi:10.1002/jcc.21707 1029 (2011). 1030 50 Kursa, M. B. & Rudnicki, W. R. Feature Selection with the Boruta Package. Journal of Statistical Software 36, 1 - 13, doi:10.18637/jss.v036.i11 (2010). 1031 1032 Ribeiro, M. T., Singh, S. & Guestrin, C. in Proceedings of the 22nd ACM SIGKDD 51 1033 International Conference on Knowledge Discovery and Data Mining 1135–1144 1034 (Association for Computing Machinery, 2016). 1035 52 Ozaydin, B., Burd, H., Lee, T. S. & Keasling, J. D. Carotenoid-based phenotypic screen 1036 of the yeast deletion collection reveals new genes with roles in isoprenoid production. 1037 Metab Eng 15, 174-183, doi:10.1016/j.ymben.2012.07.010 (2013). 1038 53 Nakayasu, E. S. et al. MPLEx: a Robust and Universal Protocol for Single-Sample Integrative Proteomic, Metabolomic, and Lipidomic Analyses. mSystems 1, 1039 1040 doi:10.1128/mSystems.00043-16 (2016). 1041 54 Stow, S. M. et al. An Interlaboratory Evaluation of Drift Tube Ion Mobility-Mass 1042 Spectrometry Collision Cross Section Measurements. Anal Chem 89, 9048-9055, 1043 doi:10.1021/acs.analchem.7b01729 (2017). 1044 Bilbao, A. et al. A Preprocessing Tool for Enhanced Ion Mobility-Mass Spectrometry-55 1045 Based Omics Workflows. J Proteome Res 21, 798-807, 1046 doi:10.1021/acs.jproteome.1c00425 (2022). 1047 56 Kall, L., Storey, J. D., MacCoss, M. J. & Noble, W. S. Posterior error probabilities and 1048 false discovery rates: two sides of the same coin. J Proteome Res 7, 40-44, 1049 doi:10.1021/pr700739d (2008). 1050 Stratton, K. G. et al. pmartR: Quality Control and Statistics for Mass Spectrometry-Based 57 Biological Data. J Proteome Res 18, 1418-1425, doi:10.1021/acs.jproteome.8b00760 1051 1052 (2019).1053 58 Webb-Robertson, B. J. et al. Combined statistical analyses of peptide intensities and 1054 peptide occurrences improves identification of significant peptides from MS-based 1055 proteomics data. J Proteome Res 9, 5748-5756, doi:10.1021/pr1005247 (2010). 1056 59 Brandl, J. et al. A community-driven reconstruction of the Aspergillus niger metabolic 1057 network. Fungal Biol Biotechnol 5, 16, doi:10.1186/s40694-018-0060-7 (2018).

1058 60 Kim, J. et al. Multi-Omics Driven Metabolic Network Reconstruction and Analysis of 1059 Lignocellulosic Carbon Utilization in Rhodosporidium toruloides. Front Bioeng Biotechnol 1060 8, 612832, doi:10.3389/fbioe.2020.612832 (2020). Nogales, J. et al. High-quality genome-scale metabolic modelling of Pseudomonas 1061 61 1062 putida highlights its broad metabolic capabilities. Environ Microbiol 22, 255-269, 1063 doi:10.1111/1462-2920.14843 (2020). 1064 62 King, Z. A. et al. Escher: A Web Application for Building, Sharing, and Embedding Data-1065 Rich Visualizations of Biological Pathways. PLoS Comput Biol 11, e1004321, 1066 doi:10.1371/journal.pcbi.1004321 (2015). 1067 Ebrahim, A., Lerman, J. A., Palsson, B. O. & Hyduke, D. R. COBRApy: COnstraints-63 Based Reconstruction and Analysis for Python. BMC Syst Biol 7, 74, doi:10.1186/1752-1068 1069 0509-7-74 (2013). 1070 64 Kim, Y. M. et al. Diel metabolomics analysis of a hot spring chlorophototrophic microbial mat leads to new hypotheses of community member metabolisms. Front Microbiol 6, 1071 1072 209, doi:10.3389/fmicb.2015.00209 (2015). 1073 65 Hiller, K. et al. MetaboliteDetector: comprehensive analysis tool for targeted and 1074 nontargeted GC/MS based metabolome analysis. Anal Chem 81, 3429-3439, 1075 doi:10.1021/ac802689c (2009). 1076 MacLean, B. et al. Skyline: an open source document editor for creating and analyzing 66 1077 targeted proteomics experiments. Bioinformatics 26, 966-968, 1078 doi:10.1093/bioinformatics/btg054 (2010). 1079 Cox, J. & Mann, M. MaxQuant enables high peptide identification rates, individualized 67 1080 p.p.b.-range mass accuracies and proteome-wide protein quantification. Nat Biotechnol 1081 26, 1367-1372, doi:10.1038/nbt.1511 (2008). Bilbao, A. PeakDecoder enables machine learning-based metabolite annotation and 1082 68 1083 accurate profiling in multidimensional mass spectrometry measurements, 2023). 1084 69 Bilbao, A. in Encyclopedia of Bioinformatics and Computational Biology (eds Shoba 1085 Ranganathan, Michael Gribskov, Kenta Nakai, & Christian Schönbach) 84-95 1086 (Academic Press, 2019). 1087

1088 Acknowledgements

1089 This work was part of the DOE Agile BioFoundry (http://agilebiofoundry.org) supported by the 1090 U.S. Department of Energy, Energy Efficiency and Renewable Energy, Bioenergy Technologies 1091 Office (to K.E.B.J.), and used capabilities developed under the National Institute of General 1092 Medical Sciences grant P41 GM103493 (to R.D.S.). Microbial biomass samples from 1093 fermentations on different hydrolysates were generated as a part of the Feedstock Conversion 1094 Interface Consortium, funded by the Department of Energy Bioenergy Technologies Office.

1095 Author contributions

K.E.B.J., A. Bilbao and N.M. planned this work. A. Bilbao conceived the PeakDecoder algorithm,
implemented the software, performed the LC-IM-MS data analyses and wrote the manuscript.
N.M. prepared the metabolomics standards and microbial samples for LC-MS acquisition,
optimized the LC method, performed the SRM and GC-MS metabolomics experiments and wrote
the interpretation of the *P. putida* and *R. toruloides* results. J.K. performed all the metabolite
clustering, pathway and statistical analyses and wrote the interpretation of the *A. pseudoterreus*

and *A. niger* results. D.J.O. optimized the method parameters and acquired the LC-IM-MS data.
Y.G. performed the SRM and global proteomics experiments and data analysis.

1104 K. Poorey and A.G. implemented the AMSS tool. K.W. provided support for optimizing the LC 1105 method and contributed to the development of the SRM metabolomics method. M.B. and C.D.N. 1106 performed sample extraction and preparation. K. Pomraning, S.D. and Z.D. planned the 1107 experimental design, engineered, and grew the A. pseudoterreus and A. niger strains. R.W. 1108 planned the experimental design, engineered, and grew the *P. putida* strains. E.O. planned the 1109 experimental design, engineered, and grew the R. toruloides strains. E.S.B. provided insightful 1110 comments about IM. C.J.P., R.A.F., A.T., and A.A. contributed to the development and 1111 optimization of the LC method and the AMSS tool. Y.K. supervised the initial LC method 1112 development and provided helpful comments about metabolomics. D.T., J.G., R.D.S., J.K. 1113 Michener, J.M.G., and J.K. Magnuson managed related projects. K.E.B.J. supervised this work 1114 and assisted in preparing the manuscript. All authors discussed, edited, and approved the final 1115 manuscript.

1116 Competing interests

A.G., R.A.F., A.T., and A.A. are employees at Agilent Technologies. The remaining authorsdeclare no competing interests.

1119 Tables

1120 Table 1. Summary of PeakDecoder training performance and identification results in 1121 microbial samples. Number of unique metabolites and number of annotated features refer to the 1122 identifications obtained by matching against our library of 64 metabolites by accurate mass plus 1123 either 2 dimensions with RT-CSS (annotations at the MS1 level only, i.e., no detected fragments) 1124 or 3 dimensions with RT-CCS-DIA (i.e., including fragments). Unique metabolites do not count 1125 repetitions. Annotated features include sample replicates and different conditions. Data for the A. 1126 pseudoterreus and A. niger samples were acquired with 20V collision energy (CE) and for P. 1127 putida and R. toruloides with both 20V and 40V CE. The number of annotations varies per dataset 1128 due to CEs and their different number of conditions and replicates.

Dataset (number of runs)	Collision energy	# Target peak- groups in training	Training accuracy (average 10-fold cross- validation)	# Unique metabolites		# Annotated features	
				RT- CCS	RT-CCS- DIA	RT-CCS	RT-CCS- DIA
A. pseudoterreus & A. niger (46)	20V	234	98.50	12	27	322	909
P. putida (22)	20V & 40V	5152	98.86	12	25	329	239
R. toruloides (48)	20V & 40V	1400	97.04	14	22	636	248

1129 1130

1131 Figure legends

1132 Figure 1. Analytical workflow for multidimensional metabolite profiling by LC-IM-MS and

1133 data structure. Metabolite extracts are separated by LC, followed by IM, and analyzed by MS 1134 in the All-Ions DIA mode which alternates between low and high collision energies to capture 1135 precursor and fragment ion spectra within the same run. Spectra are represented by gray 1136 dashed lines. Rather than collecting a single spectrum at every LC time point, coeluting ions 1137 (i.e., with close elution times) in this example at the 2nd order of elution and represented by 1138 spheres and peaks, in blue, red and orange colors, could be further distinguished by the ion 1139 mobility separation where multiple spectra are collected into IM frames. Fragments are detected within the same elution and mobility time window as their precursors. Figure adapted from 1140

- 1141 previous work⁶⁹, with permission from Elsevier.
- 1142

1143 Figure 2. Computational workflow for multidimensional metabolite profiling by LC-IM-MS. 1144 a PeakDecoder algorithm. Step-1: data is processed in untargeted mode (UFD, MS-DIAL) to 1145 extract all precursor ion features (MS1) and their respective deconvoluted fragment ions (pseudo 1146 MS2) based on co-elution and co-mobility. Step-2: a preliminary training set is generated by using 1147 the detected and deconvoluted peak-groups as targets and producing their corresponding 1148 decoys. Step-3: targeted data extraction is performed (TDX, Skyline) to extract the precursor and 1149 fragment ion signals for the training set from all the LC-IM-MS runs and export their XIC metrics. 1150 Step-4: an SVM classifier is trained using multiple scores calculated from the XIC metrics of the 1151 training set. Before training, filtering for high-quality fragments is applied to keep high-quality 1152 peak-groups as targets (i.e., based on various thresholds for metrics of precursor and at least 3 1153 fragments; details in Methods) and their corresponding decoys in the final training set. The model 1154 learns to distinguish true and false co-elution and co-mobility, independently of the features' 1155 metabolite identity. Step-5: TDX is performed to extract the signals of the guery set of metabolites 1156 in the library from all the LC-IM-MS runs and export their XIC metrics. Step-6: the trained model 1157 is used to determine the PeakDecoder score of the query set of metabolites and estimate an FDR. 1158 **b** Example of decoy generation. The detected and deconvoluted peak-groups are associated by 1159 pairs and used as targets. For each pair of targets, A and B (fragments represented in red and 1160 blue colors, respectively), a pair of decoys is generated by keeping the same precursor and its 1161 properties and swapping the m/z values of 40-60% of the fragments (from the 6 most intense in 1162 this example). XIC metrics for targets correlate well with expected values but deviations and low 1163 spectral similarity occur for decoys (examples indicated in orange).

1164

1165 Figure 3. Analysis of microbial samples by LC-IM-MS using PeakDecoder. a Comparison of 1166 scores in training. Targets and decoys are represented by blue and red colors, respectively. 1167 Distributions of LC-IM-MS peak-groups by each individual score (highlighted in orange) showed 1168 limited separation of targets and decoys. Individual scores used as machine learning features 1169 were combined into the composite PeakDecoder score providing an improved separation power 1170 and resulted in a larger number of true positives for lower FDR thresholds than the cosine 1171 similarity score, which is the best score individually. b Example of chromatograms and filtered ion 1172 mobility window. Signals for 'fructose 1,6-diphosphate (F16DP)' from the standard (precursor m/z1173 338.98877, RT 4.95 min, CCS 155.00 and 6 fragments) and corresponding peaks from a microbial

1174 sample (annotated by PeakDecoder). Chromatograms show the same relative abundances in the 1175 standard and the microbial sample confirming the correct metabolite annotation based on 1176 fragmentation pattern and RT. The IM frame at the LC apex shows the filtering window 1177 corresponding to the expected CCS and highlights the precursor with multiple isotopic peaks. c 1178 Benchmarking of identification performance compared to manual curation. True positives (TP) 1179 and false positives (FP) are represented by blue and red colors, respectively. PeakDecoder at 1180 1% estimated FDR increased TP annotations (211) compared to MS-DIAL (TP=70, total score > 1181 60) and decreased by 4 compared to Skyline (TP=215, cosine similarity > 0.8), while decreasing FP annotations (FP: PeakDecoder=4, MS-DIAL=13, Skyline=15). Results from the P. putida 1182 1183 samples (n=22). Source data are provided as a Source Data file.

1184

1185 Figure 4. Annotation selectivity by different analytical separations in microbial samples. a 1186 A. pseudoterreus and A. niger (n=46). **b** P. putida (n=22). **c** R. toruloides (n=48). Bars represent 1187 the number of possible LC-IM-MS peaks from untargeted feature detection results matched within 1188 tolerances. Colors represent the type of match: red=Mass, yellow=Mass-RT, blue=Mass-CCS, 1189 and purple=Mass-RT-CCS. In all three microbial datasets, using accurate mass alone resulted in 1190 the highest number of features, notably for the metabolites with smaller masses. Combining 1191 accurate mass to either RT or CCS reduced the number of matched features. By combining 1192 accurate mass with both RT and CCS, the number of possible features was reduced to one in 1193 most cases. These results illustrate the power of multidimensional separations to increase the 1194 annotation confidence and quantitation accuracy in metabolomics studies by resolving the high 1195 degree of structural diversity derived from isomers and isobars. Source data are provided as a 1196 Source Data file.

1197

1198 Figure 5. Metabolomics profiling of 3HP-producing A. pseudoterreus and A. niger strains. 1199 a Relative and label-free intracellular metabolites levels quantified by PeakDecoder (n=46). Red. 1200 yellow, and blue colors indicate high, medium, and low log2 intensity values, and gray color 1201 indicates missing values. b CCS errors of the good-quality features in 24 samples confirmed the 1202 detection of 3HP (green bar, 113.8 CCS) instead of lactic acid (orange bar, 113.0 CCS), which is an isomeric molecule (same formula but with different 3D structure). c Metabolites in the 3HP 1203 1204 production pathway and their log2 fold changes over the control sample (parent strain). Statistical 1205 analysis was performed using the IMD-ANOVA method. Stars indicate statistically significant 1206 changes (*: p-value < 0.05, **: p-value < 0.01 and ***: p-value < 0.001). Y-axis for pyruvic (A. 1207 pseudoterreus) and 2,4-diaminobutanoic acids represent mean log2 intensity due to no detection 1208 in the control strain. Source data are provided as a Source Data file.

1209

Figure 6. Metabolomics and proteomics profiling of *P. putida* wild type and engineered muconate-catabolizing strains. a Relative and label-free intracellular metabolites levels quantified by PeakDecoder (n=22, with 11 samples and 2 collision energies per sample). Red, yellow, and blue colors indicate high, medium, and low log2 intensity values, and gray color indicates missing values. b Glucose and muconate catabolism pathways of mucK PP5042 and fold changes compared to the wild-type strain. Circles indicate metabolites and arrows indicate proteins detected by SRM. Symbols indicate protein detection: * detected in the wild type but not

- detected in the mucK samples and # detected in the mucK but not in the wild type. Source dataare provided as a Source Data file.
- 1219

1220 Figure 7. Metabolite and enzyme levels in the mevalonate pathway of *R. toruloides* strains.

1221 **a** Relative and label-free abundance levels are represented in blue for metabolomics (n=48, with

1222 24 samples and 2 collision energies per sample) and black for proteomics (n=24 samples). Strains

were grown in hydrolysates with different contents of ash and moisture and collected at 36 and
60 hr. **b** Bisabolene production (extracellular) captured in a dodecane overlay. Data are presented

1225 as mean values with error bars from standard deviation of 3 biological replicates. Source data are

1226 provided as a Source Data file.

1227 Supplementary information

1228 Supplementary information.pdf

1229 Supplementary files

- 1230 Supplementary Data 1.zip
- 1231















Strain malic acid trehalose mannitol isocitric acid citric acid 2,4-diaminobutanoic acid succinate semialdehyde alpha-ketoglutaric acid (aKG) ribose 5-phosphate 3-hydroxypropanoic acid (3HP) pyruvic acid UDP-glucose glucose 1-phosphate (G1P) aspartic acid aspartate semialdehyde adenosine L-threonine 4-aminobutyric acid (GABA) succinic acid glucose 6-phosphate (G6P) fructose 6-phosphate (F6P) 3-phosphoglyceric acid (3PG) glutathione reduced b phenylalanine xylitol arabitol alanine tryptophan cis-aconitic acid uridine gluconic acid glutamic acid glutamine trans-aconitic acid alucose citramalic acid fumaric acid

cis,cis-muconic acid





