1	Functional screening and in vitro analysis reveals thioesterases with enhanced substrate
2	specificity profiles that improve short-chain fatty acid production in Escherichia coli
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10	Running Head: Thioesterases for short-chain fatty acid biosynthesis
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13 Abstract

Short-chain fatty acid (SCFAs) biosynthesis is pertinent to production of biofuels, industrial 14 compounds, and pharmaceuticals from renewable resources. To expand on *Escherichia coli* 15 SCFA products, we previously implemented a coenzyme A (CoA)-dependent pathway that 16 condenses acetyl-CoA to a diverse group of short-chain fatty acyl-CoAs. To increase product 17 titers and reduce premature pathway termination products, we describe in vivo and in vitro 18 analyses to understand and improve the specificity of the acyl-CoA thioesterase enzyme, which 19 releases fatty acids from CoA. A total of 62 putative bacterial thioesterases, including 23 from 20 21 the cow rumen microbiome, were inserted into a pathway that condenses acetyl-CoA to an acyl-CoA molecule derived from exogenously provided propionic or isobutyric acid. Functional 22 screening revealed thioesterases that increase production of saturated (valerate), unsaturated 23 (trans-2-pentenoate) and branched (4-methylvalerate) SCFAs compared to overexpression of E. 24 coli thioesterase tesB or native expression of endogenous thioesterases. To determine if altered 25 thioesterase acyl-CoA substrate specificity caused the increase in product titers, six of the most 26 promising enzymes were analyzed *in vitro*. Biochemical assays revealed that the most productive 27 thioesterases rely on promiscuous activity, but have greater specificity for product-associated 28 29 acyl-CoAs than for precursor acyl-CoAs. Here we introduce novel thioesterases with improved specificity for saturated, branched and unsaturated short-chain acyl-CoAs, thereby expanding the 30 diversity of potential fatty acid products while increasing titers of current products. The growing 31 32 uncertainty associated with protein database annotations denotes this study as a model for isolating functional biochemical pathway enzymes in situations where experimental evidence of 33 enzyme function is absent. 34

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36 **1. Introduction**

The potential for producing biofuels, industrial compounds, and pharmaceuticals from 37 renewable resources has led to an increased interest in short-chain (C2 - C7) fatty acid (SCFA) 38 biosynthesis. Developing recombinant strains to produce these molecules could lead to 39 production of polymers such as polyhydroxyalkanoates and pharmaceuticals such as statins from 40 fossil fuel alternatives (1, 2). Supporting the development of microbially-synthesized products is 41 the reduced cost and increased prevalence of genomic sequencing. The resulting profusion of 42 metabolic diversity provides a wealth of potential enzymes with known genetic sequences for 43 44 improving biosynthetic pathways. For example, recent genomic sequencing has unveiled the metabolic diversity of important members from the cow rumen microbiome, an environment rich 45 in SCFAs (3, 4). These genome sequences provide an opportunity to find enzymes that improve 46 production and specificity in SCFA biosynthesis pathways. 47

One such pathway that would benefit from improved enzyme specificity is Coenzyme A 48 (CoA)-dependent biosynthesis of SCFAs. In the CoA-dependent pathway, a thiolase enzyme 49 condenses an acyl-CoA molecule with acetyl-CoA (Figure 1). The resulting 3-ketoacyl-CoA 50 molecule is then sequentially reduced by reductase, crotonase, and enoyl-reductase enzymes 51 before the 3-hydroxy, unsaturated, and saturated fatty acids, respectively, are cleaved from CoA 52 using a thioesterase enzyme. Previous work from our lab has used this pathway with acetyl-CoA 53 and glycolyl, propionyl, or isobutyryl-CoA as the condensed substrates to produce a variety of 54 55 SCFAs using *E. coli* TesB including 3-hydroxyvalerate (3-hydroxypentanoate) (1), dihydroxybutyrate (5), 3-hydroxy-4-methylvalerate (5), and a variety of alcohols (6). One benefit 56 of CoA-dependent biosynthesis of SCFAs is the diversity of potential products (7); however, 57 58 with this diversity comes a need for selective enzymes that increase final product titers by

59	minimizing substrate flux to undesired byproducts. The selectivity of the final enzyme in the
60	pathway, the thioesterase, is of particular importance because it influences the product profile by
61	catalyzing fatty acid release from CoA at each step of the pathway (Figure 1) and is important
62	for secretion of fatty acid products (8). While many acyl-ACP thioesterases have been
63	investigated for improved fatty acid production, acyl-CoA thioesterases are not as well explored
64	(8-10). Despite its preference for acyl-CoAs in the C14-C18 range (11), the <i>E. coli</i> acyl-CoA
65	thioesterase TesB produces diverse SCFAs (12). However, locating more selective thioesterases
66	may reduce byproduct formation and increase final product titers.
67	Selecting individual thioesterase enzymes for functional screening against short-chain
68	fatty acyl-CoAs is challenging because much of their vast phylogenetic and functional diversity
69	is poorly understood (13). While many thioesterases have been explored for long-chain fatty acid
70	production (14-17), few studies have focused on those that prefer short-chain acyl-CoAs. Several
71	broad specificity acyl-CoA thioesterases including E. coli TesB and Saccharomyces cerevisiae
72	Pte1p can be used for SCFA production, but lack the specificity necessary for optimizing
73	biosynthetic pathways (11, 18). One approach to selecting thioesterases for functional screening
74	that improves the likelihood of finding enzymes with the desired specificity is to investigate
75	those proteins with similarity to commonly used and effective enzymes. However, the absence of
76	known selective short-chain acyl-CoA thioesterases restricts this approach. Sampling candidates
77	more broadly will increase opportunities for finding enzymes with new substrate specificities,
78	but will also increase the number of thioesterases with undesired activities. Some combination of
79	these routes can be used to screen sufficient phylogenetic breadth while also increasing the
80	sample size of TesB-like thioesterases to reveal enzymes with greater specificity for short-chain
81	acyl-CoAs.

To address this need, we functionally screened 62 putative thioesterases in a pathway for 82 production of the SCFAs 3-hydroxyvalerate, trans-2-pentenoate, valerate, and 4-methylvalerate 83 in E. coli. Six enzymes were homologous to variants active on short-chain acyl-CoAs, while the 84 remaining 56 represent all of the annotated thioesterases from five bacterial strains of interest. 85 Based upon *in vivo* fatty acid titers of those recombinant strains, six enzymes were chosen for *in* 86 vitro analysis on a broad range of acyl-CoAs to determine their substrate preferences. The 87 combination of *in vivo* and *in vitro* data indicate that we have uncovered thioesterases with 88 greater specificity for and production of unsaturated, saturated, and branched SCFAs in E. coli 89 90 relative to TesB and endogenous thioesterase activity. 91 2. Methods 92 2.1. Bacterial Strains 93 Rumen isolate Prevotella ruminicola 23 was obtained from Dr. Roderick Mackie of the 94 University of Illinois, Urbana-Champaign (USA). Genomic DNA from rumen isolates 95 Fibrobacter succinogenes S85 and Ruminococcus albus 7 were obtained from Dr. Paul Weimer 96 of the US Dairy Forage Research Center, USDA-Agricultural Research Service, Madison (USA). 97 98 Alcanivorax borkumensis SK2 (ATCC# 700651) and genomic DNA from Pseudomonas aeruginosa PAO1 (ATCC# 47085) were purchased from the ATCC. Genomic DNA of 99 Rhodopseudomonas palustris CGA009 was obtained from Dr. Caroline Harwood of the 100 101 University of Washington, Seattle (USA). Pseudomonas syringae py. maculicola ES4326 was obtained from Dr. Fred Ausubel at Massachusetts General Hospital (Boston, MA, USA). E. coli 102 103 MG1655 (DE3) Δ endA Δ recA was previously constructed in our lab (1). 104

105 2.2. Plasmid and Strain Construction

106	Plasmid pET/ter/bktB/pct was constructed previously (6) and plasmid pCDF/phaB/phaJ4
107	was constructed by subcloning <i>phaB</i> from pET/ <i>bktB</i> / <i>phaB</i> (5) into multiple cloning site I by
108	NdeI/XhoI restriction digest and <i>phaJ4</i> into multiple cloning site II of pCDFDuet-1
109	(EMDmillipore) using primers listed in Table S1. Plasmid pET/bktB/pct was constructed from
110	pET/ter/bktB/pct by BamHI/NotI restriction digest followed by treatment with Mung Bean
111	Nuclease (New England Biolabs) and blunt ligation of the 8 kb fragment.
112	Genomic DNA was isolated from bacterial strains using the Wizard Genomic DNA
113	Purification Kit (Promega). Custom oligonucleotides were purchased for the PCR amplification
114	of all individual thioesterases and CoA ligases from purified genomic DNA (Integrated DNA
115	Technologies). Primers used for amplification are listed in Table S1. Following amplification,
116	individual genes were inserted into the expression vector pACYCDuet-1 (EMDmillipore) using
117	polymerase incomplete primer extension (PIPE)-based cloning (19). The genes encoding R .
118	palustris CoA ligases FcsA (Rpa4267) and VcsA (Rpa3299) (20) and the genes encoding the six
119	thioesterases chosen for further analysis were inserted into vector pTEV5 for protein purification
120	using PIPE-based methods (21). The pTEV5 construct produced an enzyme with an N-terminal
121	hexahistidine tag removable by TEV protease. Due to solubility problems in pTEV5, the genes
122	encoding thioesterases Pr1687 and Fs2108 were amplified from genomic DNA and cloned using
123	restriction enzymes NdeI and EcoRI into pTYB22 (New England Biolabs) which produced an
124	enzyme with an N-terminal chitin-binding domain removable by intein self-cleavage. Plasmid
125	sequences were confirmed using PCR amplification and DNA sequencing (GENEWIZ).
126	Gene deletions of <i>yciA</i> , <i>yigI</i> , and <i>tesB</i> in <i>E. coli</i> MG1655 (DE3) Δ <i>endA</i> Δ <i>recA</i> were made
127	using P1 transduction with strains JW1245-1, JW5588-1, and JW0442-1, respectively, from the

Keio collection as donor cells (22). The kanamycin resistance gene was removed using FLP-mediated recombination as previously described (23).

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131 2.3. Culture Conditions

Recombinant strains of E. coli MG1655 (DE3) Δ endA Δ recA were grown at 30°C in 132 Luria-Bertani (LB) medium overnight in a shaking incubator at 250 rpm. 50 µL of the overnight 133 culture was used to inoculate a 50 mL LB culture supplemented with 10 g/L glucose containing 134 100 mg/L ampicillin, 50 mg/L streptomycin, and, when pACYCDuet-1 was present, 16 mg/L 135 chloramphenicol. Cultures were grown at 30°C until an optical density at 600 nM (OD_{600}) of 0.8 136 was reached, at which point isopropyl-β-D-1-thiogalactopyranoside (IPTG) was added to a final 137 concentration of 1 mM with either propionate or isobutyrate to a final concentration of 15 mM. 138 139 Cultures were incubated at 30°C for 48 h post-induction prior to fatty acid titer determination. 140

141 2.4. Fatty Acid Analysis

Culture samples were centrifuged to pellet cells and culture supernatant was removed for
HPLC analysis. A 5 μl sample of culture supernatant was injected into an Agilent 1100 series
instrument equipped with refractive index detection (RID). Samples were processed through an
Aminex[®] HPX-87H anion-exchange column (Bio-Rad Laboratories) with isocratic flow of 5 mM
H₂SO₄ mobile phase at a rate of 0.6 mL/min and column and detector temperatures set to 35°C.
Concentrations of valerate, 3-hydroxyvalerate, trans-2-pentenoate, and 4-methylvalerate were
determined using linear regression of external standards.

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150 2.5. Protein Purification

151 Acyl-CoA ligases FcsA and VcsA and thioesterases EcTesB, PpTesB, EcYdiI and Pr655 were overproduced using pTEV5 constructs in E. coli BL21Star (DE3) (Invitrogen). One liter of 152 cells were grown at 30°C in LB medium containing 100 mg/L ampicillin until an OD₆₀₀ of 0.5 153 was reached, at which point IPTG was added to the cultures at a concentration of 100 mg/L. Post 154 induction, the cells were grown for 15 h at 30°C, then harvested by centrifugation and 155 resuspended in 2.5x v/w Buffer A (50 mM Tris-HCl pH 8.0, 300 mM NaCl, 10% v/v glycerol). 156 Protein purification followed previously described protocols for nickel chelate chromatography 157 followed by cleavage with TEV protease (24). Proteins were flash frozen in liquid nitrogen and 158 159 the concentration was determined using the Bradford assay with bovine serum albumin as a standard (25) (Bio-Rad). 160

Pr1687 and Fs2108 protein was purified using the Intein-Mediated Purification with an Affinity Chitin-binding Tag (IMPACT) expression vector pTYB22 (New England Biolabs). Cells were grown and induced using the same conditions as described for pTEV5 constructs and proteins were purified with chitin-affinity chromatography followed by intein cleavage mediated by dithiothreitol (DTT) using previously described conditions (26). Proteins were stored and quantified as described for nickel chelate chromatography purified proteins.

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168 2.6. Enzymatic Synthesis of Acyl-CoAs

Acyl-CoA synthesis reactions were carried out in 50 mM HEPES pH 7.5, 1 mM DTT, 5

mM MgCl₂, 5 mM ATP, and 2 mM CoA. Fatty acid substrate was added at 7.5 mM for butyrate,

171 3-hydroxyvalerate, trans-2-pentenoate, valerate, 4-methylvalerate, hexanoate, octanoate,

decanoate, and dodecanoate while 2 mM fatty acid substrate was used for tetradecanoate. To

increase solubility, 1% and 3% w/v triton-X 100 was added to reactions with dodecanoate and

tetradecanoate respectively. CoA ligase enzymes were added at 500 nM for all reactions. VcsA
was added to butyrate, 3-hydroxyvalerate, valerate, 4-methylvalerate, and hexanoate reactions,
while FcsA was added to octanoate, decanoate, dodecanoate, and tetradecanoate reactions.
Reactions were run overnight at room temperature for all substrates except trans-2-pentenoate,
which was run at 30°C for 6 h because these conditions reduced the appearance of degradation
products. Acyl-CoA ligases were precipitated from reactions at 95°C for 5 min, and then
removed by centrifugation.

Acyl-CoA products were purified from substrates using an Agilent 1200 series HPLC 181 with diode array detection (DAD). A 100 µL reaction volume was injected onto an Agilent 182 Eclipse XDB-C18 column and separation achieved using a mobile phase of 50 mM ammonium 183 acetate, 0.1% m/v acetic acid (Solvent A) - 50 mM ammonium acetate, 0.1% m/v acetic acid, 184 70% v/v acetonitrile (Solvent B) gradient. The method began at 100% Solvent A from 0-5 min, 185 followed by a 0-100% gradient of solvent B from 5-50 minutes, followed by an isocratic step of 186 100% solvent B from 50-55 min. The gradient was run at a flow rate of 1 mL/min and CoA was 187 monitored by measuring absorbance at 258 nM. Fractions containing peaks corresponding to 188 acyl-CoAs were collected, flash frozen in liquid nitrogen, and lyophilized. Dried acyl-CoAs were 189 then resuspended in water and the concentration was determined by the absorbance at 258 nM 190 using the molar extinction coefficient of CoA (14,328 M⁻¹ cm⁻¹) within the linear range of 191 detection (27). 192

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194 2.7. Thioesterase Activity Assays

Thioesterase activity was measured using 5,5'-dithiobis-(2-nitrobenzoic acid) (DTNB)
which reacts with free CoA after it is released through thioesterase-mediated bond cleavage.

Reactions were carried out in 1 mM DTNB, 100 mM HEPES pH 8.0, 20 µM acyl-CoA and were
run in the linear range of each thioesterase. Substrate concentrations matched those used
previously for EcTesB analysis (11). Reaction progress was monitored through the change in
absorbance at 412 nm using the molar extinction coefficient of 5-thio-2-nitrobenzoate (14,150
M⁻¹ cm⁻¹), which is formed when DTNB reacts with free CoA (28).

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203 **3. Results**

204 3.1 In Vivo Functional Screening for Thioesterases that Alter Fatty Acid Production

205 Acyl-CoA thioesterase substrate specificity plays an important role in deciding the SCFA profile produced using a CoA-based pathway. For example, if the target product is a saturated 206 fatty acid, a thioesterase with broad substrate specificity will release fatty acids at each step in 207 208 the biosynthetic pathway, depleting substrate pools and reducing the final product titer (Figure 1). To find thioesterases that increase specificity for and production of 3-hydroxy, unsaturated, 209 saturated and branched fatty acids in the pathway, 62 putative thioesterases were chosen for 210 screening. A group of six thioesterases, three TesB and three TesB2 enzymes, were chosen that 211 had 38-50% amino acid identity to E. coli MG1655 TesB (EcTesB), which makes 3-hydroxy and 212 saturated fatty acids in vivo (1, 6, 29), and 40-41% amino acid identity to A. borkumensis TesB2 213 enzyme, which was previously described as having specificity for 3-hydroxy acyl-CoAs (30) 214 (Table 1). 215

To incorporate greater phylogenetic and functional diversity of thioesterases, the remaining 56 proteins encompassed all of the annotated thioesterases from five bacterial strains (Table 1). *E. coli* MG1655 (Accession: NC_000913.2) thioesterases were chosen with the dual purpose of identifying those that increase product titers of desired SCFAs for use in heterologous

220 pathways and those that reduce product titers and should be targeted for deletion from our host strain. The second source of thioesterases, Pseudomonas putida KT2440 (Accession: 221 NC 002947.3), was chosen for its phylogenetic similarity to *E. coli* and because it is a known 222 223 producer of polyhydroxyalkanoates, which may indicate the presence of thioesterases with specificity to 3-hydroxy acyl-CoAs (31). The remaining three organisms, Prevotella ruminicola 224 23 (Accession: NC 014033.1), Fibrobacter succinogenes S85 (Accession: NC 017448.1), and 225 Ruminococcus albus 7 (Accession: NC 014833.1), were chosen because they are prevalent in the 226 cow rumen microbiome and contribute to the high concentrations of SCFAs found there (3, 4, 227 228 32).

Each of the 62 putative thioesterases was individually overexpressed in *E. coli* containing 229 all the necessary genes for CoA-dependent biosynthesis of valerate (Figure 1). Previous work 230 231 from our lab identified Treponema denticola Ter, Megaphaera elsdenii Pct, and Cupriavidus necator (formerly R. eutropha H16) BktB, PhaB, and PhaJ4 as suitable upstream pathway 232 enzymes for valerate production (6, unpublished data). Cells were grown in LB containing 233 glucose until log phase growth, then pathway genes were induced and cultures were 234 supplemented with either propionate or isobutyrate for straight or branched SCFA biosynthesis, 235 respectively (Figure 1). LC-based analysis of the culture supernatant was used to screen for 236 enzymes that altered the fatty acid product profiles. Specifically, the resulting chromatograms 237 were examined for increases and decreases of 3-hydroxyvalerate, trans-2-pentenoate, valerate, 238 239 and 4-methylvalerate. Profiles produced by recombinant strains were compared to controls with either no recombinant thioesterase or with overproduced EcTesB, a thioesterase used previously 240 241 in our group (1, 5, 6).

242 Overproduction of greater than 20% of the thioesterases functionally screened in this study resulted in observable changes in substrate, intermediate, and product titers relative to the 243 control strains (Table S2). Four thioesterases, EcYciA, Pp1466, Pp3807, and Pp4975, were 244 associated with significant reduction in valerate and 4-methylvalerate production combined with 245 increased acetate and propionate titers (Table S2). Two thioesterases, Pr1510 and Fs368, were 246 associated with a production phenotype involving reduced titers of measured substrates, 247 intermediates, and final products combined with increased glucose consumption (Table S2). The 248 most logical explanation for this phenotype is channeling of substrates to long-chain fatty acid 249 250 biosynthesis. Because long-chain fatty acids could not be quantified with our HPLC system, additional experiments outside the scope of this work are required to determine the activity 251 profile of these enzymes. 252

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3.2 Host Strain Development and Identification of Thioesterases for In Vitro Analysis 254 From the 12 annotated E. coli thioesterases screened during this study, three were chosen 255 for deletion from the host strain. The tesB gene was deleted because its overexpression resulted 256 in increased titers of 3-hydroxyvalerate, which could be a final product, but also acts as a shunt 257 258 product in the formation of trans-2-pentenoate or valerate; yciA was deleted because its overexpression increased final titers of the precursor-derived and exogenously supplied acids 259 acetate and propionate, which correlates with previous work (33), while decreasing final product 260 261 titers of valerate and 4-methylvalerate; and *vigI* was deleted because its overexpression resulted in decreased production of 4-methylvalerate and increased final acetate and isobutyrate titers. 262 The resulting triple thioesterase deletion strain E. coli MG1655 (DE3) Δ endA Δ recA Δ tesB 263 264 $\Delta vciA \Delta vigI$ was used for further *in vivo* analysis of several active thioesterases.

265 From the full set of 62 functionally screened thioesterases, EcTesB, PpTesB, EcYdiI, Fs2108, Pr655, and Pr1687 were chosen for more detailed in vivo and in vitro analysis because 266 their overproduction resulted in increased titers of 3-hydroxyvalerate, trans-2-pentenoate, 267 valerate, or 4-methylvalerate. Propionate and isobutyrate feeding experiments were performed in 268 triplicate with these six thioesterases in both E. coli MG1655 (DE3) Δ endA Δ recA and the triple 269 thioesterase deletion strain containing pET/ter/bktB/pct and pCDF/phaB/phaJ4 in an effort to 270 distinguish recombinant thioesterase activity from background and potentially increase final 271 product titers (Figure 2a, b). The greatest 3-hydroxyvalerate titers were produced in recombinant 272 273 strains overproducing TesB enzymes from E. coli and P. putida when compared with the other 60 putative thioesterases (Figure 2). The maximum titer (2.163 g/L) resulted from 274 overproduction of EcTesB (Table 2), and is similar to previously published titers from our lab 275 276 (1). 3-hydroxyvalerate titers were 20% lower in the triple deletion strain overproducing EcTesB than in *E. coli* with native thioesterases. 277 Observable titers of unsaturated SCFAs were uncommon among the recombinant strains 278 with only Pr655 overproduction resulting in detectable trans-2-pentenoate during initial 279 screening (Figure 2a, b). To determine whether strains containing the other five thioesterases 280 281 selected for further analysis could produce detectable unsaturated SCFAs in vivo, each was

introduced into a strain lacking the downstream enoyl-reductase gene *ter*, which does not

283 produce the favored substrate valeryl-CoA (Figure 2c, d). In these recombinant strains, product

titers of trans-2-pentenoate were greatest for EcYdiI, Pr655, and Pr1687, with EcYdiI

overproduction resulting in the greatest trans-2-pentenoate titer, 695 mg/L (Table 2). To our

knowledge, this is the largest published titer of an unsaturated SCFA from an engineered

pathway in *E. coli*.

288 Three thioesterases, Fs2108, Pr1687, and EcYdiI, were chosen for further analysis because their overproduction increased titers of the saturated acids valerate and 4-methylvalerate 289 while decreasing shunt product titers (Figure 2a, b). Overproduction of thioesterase Fs2108 290 achieves the greatest final titers for both the straight SCFA valerate and the branched SCFA 4-291 methylvalerate (Table 2) with 3-hydroxy and unsaturated fatty acid intermediate titers below the 292 limit of detection. While overproduction of Fs2108 resulted in a modest 15% improvement in 293 valerate titer over the EcTesB control, an improvement of greater than 200% was observed for 4-294 methylvalerate titer over the same control. This 200% improvement was possible in part because 295 4-methylvalerate titers were significantly lower than valerate titers for all 62 thioesterases tested, 296 leaving more room for improvement. Overproduction of Fs2108 in the deletion strain resulted in 297 approximately 10% more valerate and 30% more 4-methylvalerate than in the host containing all 298 299 native thioesterases, which likely results from reduced hydrolysis of 3-hydroxyacyl-CoA and trans-2-acyl-CoA precursors. No 4-methylvalerate production was observed from the triple 300 deletion strain when isobutyrate was supplied without thioesterase overexpression, which 301 indicates that the remaining native thioesterases have poor activity on 4-methylvaleryl-CoA 302 (Figure 2a, b). 303

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To determine the substrate preferences for the selected thioesterases, all six were
overexpressed in *E. coli* BL21Star (DE3) and purified. Overexpression of *Pr1687* and *Fs2108*required the use of a vector that inserted an N-terminal chitin binding domain to obtain soluble
protein. EcTesB, PpTesB, EcYdiI, and Pr655 were soluble with an N-terminal hexahistidine tag.
Thioesterases were then tested for *in vitro* acyl-CoA hydrolysis activity on 20 µM acetyl-CoA,

3.3 Determination of In Vitro SCFA Substrate Specificity for Active Thioesterases

311 butyryl-CoA, valeryl-CoA, hexanoyl-CoA, octanoyl-CoA, decanoyl-CoA, dodecanoyl-CoA and tetradecanoyl-CoA using an Ellman's reagent-based assay described previously for determining 312 specific activity (11) (Figure 3a). EcTesB, Fs2108 and Pr1687 showed a preference for longer-313 chain acyl-CoAs. PpTesB also showed a preference for longer acyl-CoAs, but the relationship 314 was less linear than for EcTesB, Fs2108 and Pr1687. Pr655 showed a strong preference for 315 butyryl and valeryl-CoAs and activity on acyl-CoAs longer than hexanoyl-CoA was below the 316 limit of detection for this assay. EcYdiI had low activity for all substrates tested. A recent 317 publication showing that EcYdiI has strong activity on the aromatic compound 1,4-dihydroxy-2-318 319 naphthoyl-CoA provides justification for the weak activity of this enzyme on the substrates provided in this study (34). 320

Specific activities were also measured for all six thioesterases on 3-hydroxy, unsaturated, 321 saturated, and branched valeryl-CoAs (Figure 3b). For 3-hydroxyvaleryl-CoA, EcTesB and 322 PpTesB had 6-fold greater specific activity than the next best thioesterase. Both EcYdiI and 323 Pr655 showed low activity for 3-hydroxyvaleryl-CoA, while specific activity of Fs2108 and 324 Pr1687 for the same substrate was below the limit of detection. No thioesterase had specific 325 activity greater than 2 µM CoA min⁻¹ mg protein⁻¹ on trans-2-pentenoyl-CoA. Pr655, EcTesB, 326 PpTesB, and EcYdiI displayed similar specific activities for this substrate; however, the small 327 differences in specific acitivity translated into significant deviations in trans-2-pentenoate 328 product titers when the thioesterases were overproduced in recombinant strains lacking the 329 330 downstream enzyme Ter (Figure 3b). Fs2108 and Pr1687 specific activity on trans-2-pentenoyl-CoA was below the limit of detection, but *in vivo* product profiles of strains lacking *ter* reflect a 331 slight preference for 3-hydroxyvaleryl-CoA over trans-2-pentenoyl-CoA (Figure 2d). 332

All six enzymes selected for further analysis had detectable activity on valeryl-CoA; 333 however, EcTesB, PpTesB, Pr655 and Fs2108 had much greater activity relative to EcYdiI and 334 Pr1687 (Figure 3b). Activity against the 4-methylvaleryl-CoA (branched) substrate was greater 335 than or equal to activity on straight valeryl-CoA for EcTesB, PpTesB, Fs2108, and Pr1687. 336 EcYdiI and Pr655 both showed weak activity against 4-methylvaleryl-CoA. In vitro analysis of 337 338 the enzymes associated with the greatest titers of valerate revealed that Fs2108 had greater specific activity for both valeryl-CoA and 4-methylvaleryl-CoA than either Pr1687 or EcYdiI 339 (Figure 3b). Further, Fs2108 specific activity for both 3-hydroxyvaleryl-CoA and trans-2-340 341 pentenyl-CoA precursors was below the limit of detection for in vitro assays. The greater specific activity of Fs2108 on 4-methylvaleryl-CoA and valeryl-CoA combined with reduced 342 specific activity on precursor acyl-CoAs likely accounts for the increased 4-methylvalerate and 343 valerate titers found for Fs2108. 344

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346 **4. Discussion**

In this study, 62 putative thioesterases were screened for increased product titers of 3-347 hydroxy, unsaturated, saturated and branched products of interest. From these, six were chosen 348 349 for more thorough *in vivo* and *in vitro* analysis. Relating the specific activity profiles of EcTesB, PpTesB, EcYdiI, Fs2108, Pr655 and Pr1687 with their in vivo product profiles of 3-350 hydroxyvalerate, trans-2-pentenoate, valerate, or 4-methylvalerate leads to several important 351 352 conclusions. First, the enzymes with the greatest *in vivo* product titers always had lower specific activity for the associated acyl-CoA than for alternative CoA substrates. For example, Fs2108 353 has a six-fold higher specific activity for tetradecanyl-CoA than for valeryl-CoA indicating that 354 355 valeryl-CoA is not the enzyme's preferred substrate. The disparity between the specific activities

356 of Fs2108 for these two substrates indicates that our pathway relies on the promiscuous thioesterase activity of Fs2108. This is particularly evident for two thioesterases, EcYdiI and 357 Pr1697, that were associated with increased valerate production despite having much higher 358 specific activity for long-chain acyl-CoAs in the case of Pr1687, or aromatic acyl-CoAs for 359 EcYdiI (34). The fact that all six thioesterases that were investigated *in vitro* prefer alternate 360 substrates over those provided in our pathways suggests that both protein engineering and future 361 bioprospecting efforts could further improve on the short-chain fatty acyl-CoA thioesterases 362 discovered here. 363

Comparing the *in vivo* product titers with *in vitro* substrate specificities of the six chosen 364 thioesterases also suggests that after a specific activity level of 1-5 μ M CoA min⁻¹ mg protein⁻¹ is 365 reached for a given acyl-CoA, the enzyme's activity for the pathway precursor acyl-CoAs 366 367 becomes an important factor influencing final product titer, illustrating the importance of selecting pathway thioesterases with reduced activity on precursor acyl-CoAs. For example, 368 strains overproducing thioesterases Pr1687 and EcYdiI produce more valerate than those 369 overproducing EcTesB despite EcTesB having 30-fold greater specific activity for valeryl-CoA 370 than Pr1687 and EcYdiI (Figure 2b). Accounting for this difference is the observation that 371 372 EcTesB also has strong specific activity on the precursor 3-hydroxyvaleryl-CoA that translates into 3-hydroxyvalerate production while Pr1687 does not have detectable activity on 3-373 hydroxyvaleryl-CoA. Further supporting the importance of reduced specific activity on pathway 374 375 precursors is the observation that recombinant strains overproducing Pr655 produce much less valerate than strains overproducing EcYdiI and Pr1687 (Figure 2a, b) even though Pr655 376 maintains greater specific activity for valeryl-CoA than thioesterases EcYdiI and Pr1687 (Figure 377 378 3b). The increased specific activity of Pr655 on 3-hydroxyvaleryl-CoA and trans-2-pentenoyl-

379 CoA precursors relative to EcYdiI and Pr1687 suggests that the reduced valerate titer results
380 from increased precursor acyl-CoAs hydrolysis.

Confounding our conclusion that thioesterases with low specific activity on precursor 381 acyl-CoAs have improved final product titers is the observation that recombinant strains 382 overexpressing *vdiI* produce high valerate titers with trans-2-pentenoate titers below the limit of 383 detection despite EcYdiI exhibiting higher in vitro specific activity for 20 µM trans-2-pentenoyl-384 CoA than for 20 µM valeryl-CoA (Figures 2a, 3b). In our recombinant strains, EcYdiI and the 385 other screened thioesterases are competing with the downstream pathway enzyme Ter for the 386 387 substrate trans-2-pentenoyl-CoA. One plausible explanation for the absence of detectable trans-2-pentenoate from this recombinant strain is that EcYdiI has a K_m, a measure of substrate 388 affinity, for both trans-2-pentenoyl-CoA and valeryl-CoA greater than the 20 µM concentration 389 used in our *in vitro* assay, which is reasonable to assume because EcYdiI has evolved for 390 specificity towards aromatic acids (34). If Ter has a lower K_m for trans-2-pentenoyl-CoA than 391 EcYdiI, then it could reduce the intracellular concentration of trans-2-pentenoyl-CoA below 20 392 µM, causing EcYdiI activity for this compound to become physiologically irrelevant. In this 393 situation, the intracellular valeryl-CoA concentration could increase to a point where low-level 394 395 specific thioesterase activity would result in significant valerate production. Pr655, on the other hand, could have a lower K_m for both trans-2-pentenoyl-CoA and valeryl-CoA, which would 396 compete more effectively for the trans-2-pentenoyl-CoA with downstream enzyme Ter and allow 397 398 continued low level production of trans-2-pentenoate (Figure 2a). Determination of kinetic parameters for these non-cognate substrates is limited by the low concentrations of acyl-CoAs 399 obtained from enzymatic synthesis and LC-based purification. 400

401 The thioesterase substrate preferences found by our *in vitro* experiments indicate the breadth of thioesterase functional diversity screened in this study and underline the importance 402 of sampling in the selection of pathway enzymes (Figure 3a and b). The TesB enzymes of E. coli 403 and *P. putida* represent the most phylogenetically (50% amino acid identity) and functionally 404 (Figure 3a and b) similar enzymes, while the remaining four thioesterases appear both 405 phylogenetically and functionaly disparate. The TesB thioesterases showed both similar in vivo 406 product profiles and *in vitro* specific activity profiles, which suggests that future screens aimed at 407 acquiring more diverse TesB function should sample enzymes with lower similarity at the amino 408 409 acid level than the E. coli and P. putida homologs. The diversity of the remaining 60 thioesterase phenotypes is an indication of the functional diversity inherent to bacterial thioesterases and 410 suggests that many opportunities remain for isolating enzymes with improved specificity over 411 those currently used in CoA-dependent biosynthetic pathways. 412

Our study also highlights the challenges associated with enzyme selection for metabolic 413 pathways. A common route to selecting pathway enzymes is to rely on proposed function of 414 known enzymes in databases and the literature or choose enzymes homologous to those with 415 proposed functions. Unfortunately, enzyme annotation in public databases has degraded as 416 417 functional analysis has not kept up with the rate of sequence deposition (35). The absence of credible studies on short-chain acyl-CoA thioesterases prompted us to implement a broad 418 functional screen of diverse candidates. Our results demonstrate the power of a well-designed 419 420 screen for isolating uncharacterized or poorly characterized enzymes that improve product titers. Our methods for screening and *in vitro* characterization of substrate specificity serve as a 421 template for investigating other poorly characterized enzyme functions for pathway 422 423 development. The *E. coli* thioesterase EcYdiI serves as an example of an enzyme whose

documented function may eliminate it from contention (34), but once included in our functional
screen, maintains the appropriate levels of promiscuous activity in intracellular conditions to
provide the highest specificity for unsaturated SCFAs discovered so far. It is unlikely that
selecting enzymes based solely upon annotated function would have identified this level of
activity because short-chain acyl-coA thioesterase activity remains poorly characterized.

During this study we isolated acyl-CoA thioesterases from diverse bacterial sources that 429 increase production of saturated, unsaturated, and branched SCFAs through improvements in 430 acyl-CoA substrate specificity. By comparing in vivo product profiles with in vitro specific 431 432 activities of thioesterases that produced the greatest 3-hydroxyvalerate, trans-2-pentenoate, valerate, and 4-methylvalerate titers, we discovered that the most productive thioesterases found 433 during functional screening: 1) use promiscuous activity to produce the SCFAs monitored in this 434 study and 2) maintain low specific activity for pathway precursors relative to the preferred acyl-435 CoA. These findings indicate this study as a model for isolating enzymes for biochemical 436 pathway functions that are poorly characterized in the literature. Further, the thioesterases we 437 identified provide opportunities for increasing titers of desirable products as well as developing 438 new pathways for the production of unsaturated SCFAs. We have also provided important in 439 vivo and in vitro data on the production phenotypes and substrate specificities of poorly 440 characterized acyl-CoA thioesterases for short-chain acyl-CoAs that are valuable for future 441 bioprospecting and engineering studies. 442

443

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563	Figure 2. In vivo fatty acid production. Titers of fatty acids produced from (a,c) E. coli MG1655
564	(DE3) Δ endA Δ recA and (b,d) E. coli MG1655 (DE3) Δ endA Δ recA Δ tesB Δ yciA Δ yigI
565	expressing CoA-based pathways for the biosynthesis of (a,b) valerate or 4-methylvalerate, (c,d)
566	trans-2-pentenoate, with six different thioesterases. The product titers shown were obtained from
567	three separate shake flasks for each thioesterase with error bars representing standard deviations.
568	Each strain contained plasmids pCDF/phaB/phaJ4 with either (a,b) pET/ter/bktB/pct or (c,d)
569	pET/bktB/pct. Each strain also contained plasmid pACYCDuet-1 with the thioesterases displayed
570	on the X-axis. Absence of product bars for a thioesterase indicates that titers were below the
571	limit of detection for the HPLC RID detector.
572	*4-methylvalerate titers were determined using three separate cultures for each thioesterase fed
573	15 mM isobutyrate in the place of propionate.
574	
575	Figure 3. Acyl-CoA specificity of six active thioesterases. (a) In vitro specific activity of six
576	thioesterases on C2-C14 acyl-CoAs. Specific activities represent the average from three
577	enzymatic reactions with error bars representing standard deviations. They were determined in
578	the linear range for each thioesterase with acyl-CoA concentrations of 20 μ M for C2, acetyl-
579	CoA; C4, butyryl-CoA; C5, valeryl-CoA; C6, hexanoyl-CoA; C8, octanoyl-CoA; C10, decanoyl-
580	CoA; C12, dodecanoyl-CoA; C14, tetradecanoyl-CoA. (b) In vitro specific activity of six
581	thioesterases on saturated, valeryl-CoA; branched, 4-methylvaleryl-CoA; 3-hydroxy, 3-
582	hydroxyvaleryl-CoA; unsaturated, trans-2-pentenoyl-CoA. Inset is enlarged activities of EcYdiI
583	and Pr655 for comparison of low level activities. Specific activities were determined using the

same conditions described in (a). The absence of a bar for a given substrate/thioesterase specific
activity indicates that the activity was below the limit of detection at absorbance of 412 nm.









b.

3-Hydroxypentanoate Trans-2-pentenoate



a.



b.



Table 1. Thioesterases screened for activity in a biosynthetic pathway for the production of short-chain fatty acids

TesB & TesB2 enzymes	<i>E. coli</i> MG1655	P. putida KT2440	P. ruminicola 23	R. albus 7	F. succinogenes S85
E. coli TesB ^a	EcTesA	Pp244	Pr655	Ral865	Fsu266
P. putida TesB	EcPaaI	Pp254	Pr1075	Ral874	Fsu270
R. opacus TesB	EcEntH	Pp262	Pr1498	Ral880	Fsu368
P. syringae TesB2	EcFadM	Pp301	Pr1510	Ral1843	Fsu803
A. borkumensis TesB2 ^b	EcYbfF	Pp580	Pr1668	Ral1929	Fsu944
P. aeruginoasa PAOI TesB2	EcYbgC	Pp1218	Pr1687	Ral1938	Fsu1747
	EcYbhC	Pp1466	Pr2385	Ral2059	Fsu2108
	EcYciA	Pp1980		Ral2801	
	EcYiiD	Pp2050		Ral3109	
	EcYigI	Pp2308			
	EcYdiI	Pp2318			
		Pp3281			
		Pp3807			
		Pp4105			
		Pp4180			
		Pp4181			
		Pp4975			
		Pp5198			
		Pp5331			
		Pp5356			
		PpPhaJ1			

^a(11) ^b(30)

Table 2. Maximum fatty acid titers observed for each product of the CoA-dependent pathway

	Maximum Titers (g/L)					
Strains	3-Hydroxyvalerate ^a	Trans-2-pentenoate b	Valerate ^c	4-Methylvalerate ^c		
MG1655 Δ endA Δ recA	2.163±0.078	0.695 ± 0.002	1.425 ± 0.011	0.428±0.019		
$MG1655 \Delta endA \Delta recA \Delta tesB \Delta yciA \Delta yigI$	1.940±0.099	0.609±0.015	1.540±0.034	0.570±0.041		

^aThioesterase EcTesB ^bThioesterase EcYdiI ^cThioesterase Fs2108