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Original Research Article

Isopentenyl diphosphate (IPP)-bypass mevalonate pathways for isopentenol production

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ARTICLE INFO

Article history: Received 8 September 2015 Received in revised form 2 November 2015 Accepted 7 December 2015 Available online 17 December 2015

Keywords: Isopentenol Isoprenol Mevalonate pathway Biofuel Phosphomevalonate decarboxylase IPP Toxicity Aeration

ABSTRACT

Branched C₅ alcohols are promising biofuels with favorable combustion properties. A mevalonate (MVA)based isoprenoid biosynthetic pathway for C_5 alcohols was constructed in *Escherichia coli* using genes from several organisms, and the pathway was optimized to achieve over 50% theoretical yield. Although the MVA pathway is energetically less efficient than the native methylerythritol 4-phosphate (MEP) pathway, implementing the MVA pathway in bacterial hosts such as E. coli is advantageous due to its lack of endogenous regulation. The MVA and MEP pathways intersect at isopentenyl diphosphate (IPP), the direct precursor to isoprenoid-derived C_5 alcohols and initial precursor to longer chain terpenes, which makes independent regulation of the pathways difficult. In pursuit of the complete "decoupling" of the MVA pathway from native cellular regulation, we designed novel IPP-bypass MVA pathways for C₅ alcohol production by utilizing promiscuous activities of two enzymes, phosphomevalonate decarboxylase (PMD) and an E. coli-endogenous phosphatase (AphA). These bypass pathways have reduced energetic requirements, are further decoupled from intrinsic regulation, and are free from IPP-related toxicity. In addition to these benefits, we demonstrate that reduced aeration rate has less impact on the bypass pathway than the original MVA pathway. Finally, we showed that performance of the bypass pathway was primarily determined by the activity of PMD. We designed PMD mutants with improved activity and demonstrated titer increases in the mutant strains. These modified pathways would be a good platform for industrial production of isopentenol and related chemicals such as isoprene.

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1. Introduction

Isopentenol (3-methyl-3-buten-1-ol) is a potential biofuel and important precursor for flavor compounds (prenols and isoamyl alcohol esters) and industrial chemicals such as isoprene (Chou and Keasling, 2012; Peralta-Yahya et al., 2012). Two classes of metabolic pathways have been engineered to produce isopentenol in microbial hosts: amino acid production pathways utilizing 2keto-acid intermediates (Connor and Liao, 2008; Connor et al., 2010), and isoprenoid biosynthesis pathways, including both the mevalonate (MVA) (Chou and Keasling, 2012; Withers et al., 2007; George et al., 2014; Zheng et al., 2013) and non-mevalonate pathway (methylerythritol 4-phosphate (MEP) or 1-deoxy-D- logous MVA pathway was constructed to produce isopentenol in *Escherichia coli* by expressing 7 genes (Fig. 1- Pathway O) (Chou and Keasling, 2012). To produce isopentenol, IPP is hydrolyzed by phosphatases such as NudF from *Bacillus subtilis* or NudB from *E. coli*. Although the initial performance of this pathway was low (8.3% of pathway-dependent theoretical yield), subsequent optimization has significantly improved yields and titers (George et al., 2014; Zheng et al., 2013). Most recently, isopentenol was produced at a titer of 2.2 g/L from 10 g/L glucose, which is almost 70% of apparent theoretical yield (George et al., 2015).

xylulose 5-phosphate (DXP) pathway) (Liu et al., 2013). A hetero-

A variety of engineering strategies have been applied to optimize the heterologous MVA pathway and improve isoprenoid production in *E. coli* (Pfleger et al., 2006; Anthony et al., 2009; Dueber et al., 2009; Redding-Johanson et al., 2011; Dahl et al., 2013; Zhu et al., 2014). In each case, balanced expression of pathway enzymes was required to maximize flux towards final

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http://dx.doi.org/10.1016/j.ymben.2015.12.002

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Fig. 1. Original and two modified mevalonate pathways for isopentenol production. The original mevalonate pathway (pathway O) produces isopentenyl diphosphate (IPP), which is dephosphorylated by NudB, as an intermediate. Two modified pathways were proposed in this study: direct decarboxylation of mevalonate (pathway I) or decarboxylation of mevalonate diphosphate (pathway II) followed by de-phosphorylation of isopentenyl monophosphate (IP). Numbers of ATP and enzymes required for each pathway are summarized in the table. Ac-CoA, acetyl-CoA; AAc-CoA, acetoacetyl-CoA; HMG-CoA, 3-hydroxy-3-methyl-glutaryl-CoA; PMK, phosphomevalonate kinase; PMD, phosphomevalonate decarboxylase).

products while minimizing the accumulation of toxic intermediates such as farnesyl diphosphate (FPP) (Martin et al., 2003), IPP (Withers et al., 2007; George et al., 2014; Zheng et al., 2013; Martin et al., 2003), and 3-hydroxy-3-methyl-glutaryl-CoA (HMG-CoA) (Pitera et al., 2007). For isopentenol production, the careful management of IPP levels is critical: engineering strategies to address its accumulation have included the deliberate "tuning" of the upstream MVA pathway (George et al., 2014) and the extensive overexpression of NudB, the phosphatase required to transform IPP into isopentenol (George et al., 2015).

Although the mechanism of IPP toxicity is unknown, the deleterious effects of its accumulation are clear. First, it has been demonstrated in various studies that accumulation of IPP inhibits cell growth (Withers et al., 2007; George et al., 2014; Martin et al., 2003), which prevents a bioprocess from achieving enough cell biomass to maximize product titer. Even prior to affecting cell growth, it is likely that the transient accumulation of IPP induces a variety of stress responses as has previously been observed during the accumulation of FPP (Dahl et al., 2013). Responses to both generalized (e.g., RpoS-induced (Hengge, 2008)) and conditionspecific stress (e.g., acid stress (Sun et al., 2011), oxidative stress (Adolfsen and Brynildsen, 2015) and osmotic stress (Cohen, 2014)) result in the recruitment of ATP-dependent defense mechanisms including DNA repair (Sun et al., 2011; Adolfsen and Brynildsen, 2015), ATPases (Sun et al., 2011), and ABC transporters (Cohen, 2014). The ATP cost of these processes may serve to compete with the energetically-expensive MVA pathway, reducing the yield and productivity of isoprenoid production. In the case of isopentenol production, high flux to IPP has an additional detrimental impact: through the action of E. coli native IPP isomerase (Idi), IPP can be diverted by native isoprenoid pathways that produce C₁₀- and C₁₅prenyl diphosphates (i.e. geranyl diphosphate (GPP) and FPP). The production of GPP and FPP decreases the carbon utilization efficiency of isopentenol production, and potentially inhibits MK activity, which in turn reduces MVA flux to the downstream enzyme reactions (Ma et al., 2011). Moreover, isopentenol production via IPP requires the energetically expensive ATPconsuming formation of diphosphate prior to enzymatic hydrolysis. This diphosphate formation and subsequent hydrolysis is considerably inefficient in terms of atom and energy economy. Due to these factors, the "IPP-dependency" of the MVA pathway may intrinsically limit the engineering of the MVA pathway for more efficient isopentenol production.

In this work, we successfully decouple isopentenol production from IPP formation by constructing two novel "IPP-bypass" pathways. These two IPP-bypass pathways rely on decarboxylation of either MVA or MVA monophosphate (MVAP) for isopentenol production and do not produce IPP as an essential precursor for isopentenol. These optimized pathways eliminate the negative effects of IPP accumulation such as growth inhibition, energyconsuming stress responses, diverted carbon flux, and regulatory inhibition on mevalonate kinase (MK). We envision that these two IPP-bypass pathways could open a new dimension of engineering the MVA pathway to produce isopentenol and isopentenol-derived valuable compounds such as isoprene.

2. Materials and methods

2.1. Strains and plasmid construction

All strains and plasmids used in this study are listed in Table 1. Throughout the studies, *E. coli* BW25113 strain was used for isopentenol production, and *E. coli* DH10B was used for genetic cloning. The original sequence of PMD_{hv} was obtained from NCBI database (HVO_1412, NC_013967.1), codon-optimized for expression in *E. coli* by GenScript (New Jersey, USA), and the optimized sequence was synthesized by IDT (Iowa, USA). A plasmid coding PMD_{se} was received from Dr. Miziorko at University of Missouri (Barta et al., 2012), and the coding sequence was amplified by PCR for sub-cloning to expression vectors.

2.2. Protein expression and purification

A plasmid encoding a wild type mevalonate diphosphate decarboxylase from Saccharomyces cerevisiae (PMD_{sc}) with N-terminal His-tag (pSKB3-PMD_{sc}) was transformed into *E. coli* BL21 (DE3). A seed culture of BL21 (DE3) harboring pSKB3-PMDsc was prepared by inoculating a single colony and growing it overnight in Luria-Bertani (LB) medium containing kanamycin (50 µg/mL). The seed culture was diluted in Terrific Broth supplemented with 2% glycerol and 50 µg/mL kanamycin and incubated at 37 °C until the optical density of the culture at 600 nm (OD₆₀₀) reached to 0.6–0.8. The cell culture was supplemented with isopropyl- β -D-thiogalactopyranoside (IPTG) to the final concentration of 0.5 mM and transferred to 18 °C for protein expression overnight. Cells were collected by centrifugation and resuspended in 50 mM Tris-HCl (pH 7.5) buffer containing 300 mM NaCl and 10 mM imidazole. Cells were lysed by sonication and purified by HisPur Cobalt Resins (Thermo Scientific, USA). The purified PMD_{sc} was desalted in 10 mM Tris-HCl (pH 7.5) containing 50 mM NaCl, 0.5 mM dithiothreitol (DTT) and 20% glycerol, and flash-frozen in liquid nitrogen for storage at -80°C. All PMD_{sc} mutants, PMD_{se}, and NudB were purified as described above, except that NudB was desalted in 50 mM Tris-HCl (pH 8.0) buffer containing 0.1 mM EDTA, 1 mM DTT and 20% glycerol.

2.3. Enzyme characterization and kinetics

In vitro enzyme kinetics of decarboxylases were performed as described in previous studies (Barta et al., 2012; Vannice et al., 2014). Briefly, enzymatic activity of decarboxylase was determined by a spectrophotometer assay quantifying ADP product formation, which was coupled to NADH oxidation by pyruvate kinase/lactate dehydrogenase. Assay mixtures were prepared in 50 mM HEPES-KOH (pH 7.5) containing 10 mM MgCl₂, 400 μ M phosphoenolpyruvate, 200 μ M NADH, 4 mM ATP, and 25 U of pyruvate kinase/lactate dehydrogenase (Sigma, P0294). The reaction was initiated by addition of various concentrations of MVAP from 100 μ M to 4,000 μ M, and the reaction velocity was determined by monitoring OD at 340 nm in Spectramax 384plus microplate reader (Molecular Devices, USA).

 Table 1

 List of strains and plasmids used in this study.

Strains	Description	Reference	
∆aphA	E. coli K12 BW25113 ∆aphA	Keio Collection (Baba	
∆agp	E. coli K12 BW25113 ∆agp	Keio Collection (Baba et al. 2006)	
$\Delta y q a B$	E. coli K12 BW25113 ∆yqaB	Keio Collection (Baba et al., 2006)	
ARK1a	JBEI-12056+JBEI-9348	This study	
ARK1b	JBEI-6824+JBEI-9348	This study	
ARK1c	JBEI-6831+JBEI-9348	This study	
ARK1d	JBEI-7575+JBEI-9348	This study	
ARK1e	JBEI-6818+JBEI-6833	This study	
ARK1f	JBEI-6818+JBEI-6834	This study	
ARK2a	JBEI-9310+JBEI-9314	This study	
ARK2b	JBEI-9309+JBEI-9314	This study	
ARK2c	JBEI-9312+JBEI-9314	This study	
ARK2d	JBEI-9311 + JBEI-9314	This study	
ARK2e	JBEI-12051 + JBEI-9314	This study	
ARK2f	JBEI-12051 + JBEI-12064	This study	
ARK2aa	JBEI-12050+JBEI-9314	This study	
ARK2a _{M1}	JBEI-9310+JBEI-12060	This study	
ARKZd _{M2}	JBEI-9310+JBEI-12061	This study	
ARKZd _{M3}	JDEI-9510+JDEI-12002	This study	
ARK3a	BEI-3100 + BEI-3277	This study	
ARKJD	BEI-9310 + BEI-12054	This study	
ARK5	IBEI-9310 + IBEI-12054 IBEI-9310 + IBEI-12059	This study	
Tuddo		This study	
Plasmids	Description	Reference	
IBEI-6818	pBbA5c-MevTo-MKco-PMKco	(George et al., 2014)	
IBEI-6824	pBbA5c-MevTco-BBa1002-pTrc-MKco-	(George et al., 2014)	
5	РМКсо		
JBEI-6831	pBbA5c-MTSA-BBa1002-pTrc-MKco- PMKco	(George et al., 2014)	
JBEI-6833	pTrc99a-NudB-PMDsc	(George et al., 2014)	
JBEI-6834	pTrc99a-NudB-PMDsc-Mkco	(George et al., 2014)	
JBEI-7575	pBbA5c-MTDA-BBa1002-pTrc-MKco-	Cift from Eunmi Vim	
IBEI-9309	РМКсо	Gift Holli Eulinii Kim	
5	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco	This study	
JBEI-9310	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco pBbA5c-MevTo-BBa1002-pTrc-MKco	This study This study	
JBEI-9310 JBEI-9311	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco pBbA5c-MevTo-BBa1002-pTrc-MKco pBbA5c-MTDA-BBa1002-pTrc-MKco	This study This study This study	
JBEI-9310 JBEI-9311 JBEI-9312	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco pBbA5c-MevTo-BBa1002-pTrc-MKco pBbA5c-MTDA-BBa1002-pTrc-MKco pBbA5c-MTSA-BBa1002-pTrc-MKco	This study This study This study This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco pBbA5c-MevTo-BBa1002-pTrc-MKco pBbA5c-MTDA-BBa1002-pTrc-MKco pBbA5c-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc	This study This study This study This study This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314 JBEI-9348	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco pBbA5c-MevTo-BBa1002-pTrc-MKco pBbA5c-MTDA-BBa1002-pTrc-MKco pBbA5c-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB	This study This study This study This study This study This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314 JBEI-9348 JBEI-12050	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco pBbA5c-MevTo-BBa1002-pTrc-MKco pBbA5c-MTDA-BBa1002-pTrc-MKco pBbA5c-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbA5c-MevTo-BBa1002-pTrc-MKco- aphA	This study This study This study This study This study This study This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314 JBEI-9348 JBEI-12050 JBEI-12051	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco pBbA5c-MevTo-BBa1002-pTrc-MKco pBbA5c-MTDA-BBa1002-pTrc-MKco pBbA5c-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbA5c-MevTo-BBa1002-pTrc-MKco- aphA pBbA5c-MevTo-MKco	This study This study This study This study This study This study This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314 JBEI-9348 JBEI-12050 JBEI-12051 JBEI-12052	PMKco pBbA5c-MevTco-BBa1002-pTrc-MKco pBbA5c-MevTo-BBa1002-pTrc-MKco pBbA5c-MTDA-BBa1002-pTrc-MKco pBbA5c-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbA5c-MevTo-BBa1002-pTrc-MKco- aphA pBbA5c-MevTo-MKco pSKB3-PMDsc	This study This study This study This study This study This study This study This study This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314 JBEI-9348 JBEI-12050 JBEI-12051 JBEI-12052 JBEI-12053	PMKco pBbASc-MevTco-BBa1002-pTrc-MKco pBbASc-MTDA-BBa1002-pTrc-MKco pBbASc-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-MKco pSKB3-PMDsc pSKB3-PMDsc	This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314 JBEI-9348 JBEI-12050 JBEI-12051 JBEI-12052 JBEI-12053 JBEI-12054	PMKco pBbASc-MevTco-BBa1002-pTrc-MKco pBbASc-MTDA-BBa1002-pTrc-MKco pBbASc-MTSA-BBa1002-pTrc-MKco pBbASc-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-MKco pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc	This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314 JBEI-9348 JBEI-12050 JBEI-12051 JBEI-12052 JBEI-12053 JBEI-12054 JBEI-12054	PMKco pBbASc-MevTco-BBa1002-pTrc-MKco pBbASc-MrVTo-BBa1002-pTrc-MKco pBbASc-MTDA-BBa1002-pTrc-MKco pBbASc-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-MKco pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSbA5-MevTo-DB-1002 a Tay MKab	This study This study	
JBEI-9310 JBEI-9311 JBEI-9311 JBEI-9314 JBEI-9314 JBEI-12050 JBEI-12052 JBEI-12053 JBEI-12054 JBEI-12055 JBEI-12056	PMKco pBbASc-MevTco-BBa1002-pTrc-MKco pBbASc-MevTo-BBa1002-pTrc-MKco pBbASc-MTDA-BBa1002-pTrc-MKco pBbASc-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-MKco pSKB3-PMDsc_K22M pTrc99a-PMDse pSKB3-PMDsc_T209D pBbASc-MevTo-BBa1002-pTrc-MKco- PMKco	This study This study	
JBEI-9310 JBEI-9311 JBEI-9311 JBEI-9314 JBEI-9348 JBEI-12050 JBEI-12052 JBEI-12052 JBEI-12054 JBEI-12055 JBEI-12056	PMKco pBbASc-MevTco-BBa1002-pTrc-MKco pBbASc-MevTco-BBa1002-pTrc-MKco pBbASc-MTDA-BBa1002-pTrc-MKco pBbASc-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-MKco pSKB3-PMDsc	This study This study	
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JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9314 JBEI-9348 JBEI-12050 JBEI-12051 JBEI-12053 JBEI-12054 JBEI-12055 JBEI-12056 JBEI-12056 JBEI-12059 JBEI-12059 JBEI-12060 JBEI-12061	PMKco pBbASc-MevTco-BBa1002-pTrc-MKco pBbASc-MTDA-BBa1002-pTrc-MKco pBbASc-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-BBa1002-pTrc-MKco- pKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pTrc99a-PMDsc pTrc94a-PMDsc pTrc94	This study This study	
JBEI-9310 JBEI-9311 JBEI-9312 JBEI-9312 JBEI-9314 JBEI-12050 JBEI-12051 JBEI-12052 JBEI-12053 JBEI-12055 JBEI-12056 JBEI-12057 JBEI-12058 JBEI-12059 JBEI-12060 JBEI-12062	PMKco pBbASc-MevTco-BBa1002-pTrc-MKco pBbASc-MrVa-BBa1002-pTrc-MKco pBbASc-MTDA-BBa1002-pTrc-MKco pBbASc-MTSA-BBa1002-pTrc-MKco pTrc99a-PMDsc pTrc99a-PMDsc-NudB pBbASc-MevTo-BBa1002-pTrc-MKco- aphA pBbASc-MevTo-BBa1002-pTrc-MKco- pKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pSKB3-PMDsc pTrc99a-PMDsc l145F pTrc99a-PMDsc pTr	This study This study	

2.4. Isopentenol production in E. coli

E. coli BW25113 harboring two plasmids was used for isopentenol production. Seed cultures of all production strains were prepared by growing single colonies in LB medium containing 100 μ g/mL ampicillin and 30 μ g/mL chloramphenicol overnight at 37 °C with shaking at 200 rpm. The seed cultures were diluted in EZ-Rich defined medium (Teknova, USA) containing 10 g/L glucose

(1%, w/v), 100 µg/mL ampicillin and 30 µg/mL chloramphenicol. The *E. coli* cell cultures were incubated in rotary shakers (200 rpm) at 37 °C, and 0.5 mM IPTG was added to induce protein expression at OD_{600} of 0.6–0.8. To provide different levels of aeration, identical volumes of the cell culture were split into two flasks for incubation at 30 °C with shaking at either 200 rpm or 30 rpm.

For isopentenol quantification, $250 \ \mu$ L of cell culture was combined with $250 \ \mu$ L of ethyl acetate containing 1-butanol ($30 \ mg/L$) as an internal standard. This mixture of ethyl acetate and cell culture was vigorously shaken for 15 min and subsequently centrifuged at 13,000g for 2 min to separate ethyl acetate from the aqueous phase. 100 \mu L of the ethyl acetate layer was diluted 5-fold, and 1 \mu L was analyzed by Agilent GCMS equipped with Cyclosil-B column (Agilent, USA) or Thermo GCFID equipped with DB-WAX column (Agilent, USA) for quantitation of isopentenol.

2.5. Phosphatase screening

To identify IP-hydrolyzing endogenous phosphatases, single gene knockout mutants of 36 phosphatases, of which substrates are mostly mono-phosphorylated metabolites, were retrieved from the Keio collection (Baba et al., 2006). 1 mL overnight cultures from each mutant were concentrated in 0.5 mL of 50 mM Tris–HCl (pH 8.0) buffer containing 1 mM DTT and ~50 mg of glass beads (< 100 μ m, Sigma-Aldrich, USA). Cells were lysed by bead-beating for 2 min at 6.0 M/s (MP biomedicals Fast Prep, USA). After centrifugation of cell lysates at 20,000g for 10 min, clear supernatant was used for assay reaction containing 0.5 mM isopentenyl monophosphate (IP). An equal volume of ethyl acetate was added to 100 μ L of assay reaction after incubating overnight at 30 °C, and isopentenol was extracted for 10 min by vigorous mixing.

Coding sequences of *agp*, *aphA*, and *yqaB* were amplified from BW25113 genome by PCR, and they were subsequently cloned to pBbE1a vector (Lee et al., 2011) for over-expression. All primers used in this study are listed in Supplementary Table 1. Expression of these three genes were induced by addition of 0.5 mM IPTG to the cell cultures, and cell lysates of each sample were prepared with three biological replicates as described above for screening of the 36 mutants. 600 μ L of assay reactions containing cell lysates, 1 mM DTT were prepared and the reaction was initiated by addition of 0.5 mM IP. At each time point (0, 1, 3, 6 and 22 h), 100 μ L of ethyl acetate to extract isopentenol.

2.6. Quantification of metabolites

All metabolites were analyzed by liquid chromatography mass spectrometry (LC-MS; Agilent Technologies 1200 Series HPLC system and Agilent Technologies 6210 time-of-flight mass spectrometer) on a ZIC-HILIC column (150 mm length, 2.1-mm internal diameter, and 3.5- μ m particle size). Standard chemicals (IPP and IP) were purchased from Sigma-Aldrich (USA). Metabolites were eluted isocratically with a mobile phase composition of 64% (v/v) acetonitrile containing 50 mM ammonium acetate with a flow rate of 0.15 mL/min. IPP and IP from *E. coli* extracts or enzyme assay were quantified via eight-point calibration curves ranging from 781.25 nM to 200 μ M.

3. Results and discussions

3.1. Design rationale for IPP-bypass isopentenol pathways

The biosynthesis of IPP from MVA consists of three energyconsuming reactions: two kinases (MK and phosphomevalonate



Fig. 2. GC/MS chromatogram (left) and mass spectra (right) of ethyl acetate-extracted metabolites detected from (A) control strain with three genes (ARK3a; *atoB*, *HMGS*, *HMGR*) and (B) engineered strains with four genes (ARK3b; *atoB*, *HMGS*, *HMGR* and *PMD_{sc}*). The mass spectrum of the peak that eluted at 9.49 min detected in ARK3b (B) is very similar to that of the isopentenol standard (C), and is not present in the ethyl acetate blank (D). Arrows indicate masses of the peak at retention time of 9.49 min detected from both standard (C) and the engineered strain (B; ARK3b).

kinase (PMK)) result in the formation of diphosphomevalonate (MVAPP), which is subsequently transformed by a decarboxylase (PMD) to form IPP. The diphosphate group of IPP is essential in chain elongation to produce GPP and FPP, and in the carbocation formation to produce cyclic terpenes since the removal of the diphosphate group is thermodynamically-favorable (Degenhardt et al., 2009). In isopentenol production via the MVA pathway, the alcohol is also produced by removal of the diphosphate group of IPP. However, this reaction is different from carbocation formation and does not require the diphosphate group as an essential leaving group to drive the hydrolysis reaction. Therefore, formation of the diphosphate group and its subsequent removal make the overall MVA pathway for isopentenol inefficient by unnecessarily consuming two ATPs.

To address the energetic limitations of IPP formation-and the deleterious effects of its accumulation-we designed two modified isopentenol pathways that bypass the formation of IPP (Fig. 1). The first modified pathway (pathway I) is designed for the direct conversion of MVA to isopentenol via ATP-driven decarboxylative elimination, and the second pathway (pathway II) is designed for a decarboxylative elimination of MVAP to IP followed by the hydrolysis of IP to isopentenol (Fig. 1). These modified pathways result in IPP-independent isopentenol production, which could relieve toxicity and prevent the loss of IPP flux to native pathways such as ubiquinone biosynthesis. Moreover, these two pathways reduce the complexity and energy cost of isopentenol production. As shown in Fig. 1, direct decarboxylation of MVA (pathway I) reduces the number of enzymes required from 7 to 4 and the ATP requirement per molecule of isopentenol from 3 to 1. In IPP-bypass pathway II, the number of enzymes is reduced from 7 to 5 and ATP molecules from 3 to 2 compared to the original pathway. Given the potential benefits of pathways I and II over the original MVA pathway (pathway O), we explored options to construct and express these optimized pathways in E. coli to produce isopentenol.

3.2. Engineering of IPP-bypass pathway I and identification of promiscuous decarboxylase activity toward MVA and MVAP

Engineering IPP-bypass pathways I and II requires a decarboxylase that converts MVA or MVAP to isopentenol or IP, respectively. Based on the chemical structures of the substrates and products (Fig. 1) and proposed mechanism of the decarboxylation reaction, we hypothesized that PMD might serve as a decarboxylase for MVA and MVAP in addition to its native substrate, MVAPP. Since PMD from *S. cerevisiae* (PMD_{sc}) has been widely used for isoprenoid production in engineered *E. coli* (George et al., 2014; Zheng et al., 2013; Alonso-Gutierrez et al., 2013; Kim et al., 2015; Wang et al., 2010), we initially chose PMD_{sc} as the target PMD enzyme for each bypass pathway. PMD_{sc} was previously reported to convert 3hydroxy-3-methylbutyrate (3-HMB) to isobutene (Gogerty and Bobik, 2010), which supports the hypothesis that this enzyme has promiscuous decarboxylase activity.

With PMD_{sc} as a potential decarboxylase for MVA, IPP-bypass pathway I was first constructed in *E. coli* by expressing three enzymes (AtoB, HMGS, and HMGR) to produce MVA along with PMD_{sc} (strain ARK3a, Table 1). When the strain was tested *in vivo*, the engineered *E. coli* produced 0.85 ± 0.18 mg/L isopentenol (Fig. 2B) while the control strain, which expressed only AtoB, HMGS, and HMGR without PMD_{sc} (strain ARK3b, Table 1) did not show any detectable level of isopentenol (Fig. 2A). The fragmentation pattern and retention time of the isopentenol peak detected in strain ARK3a matched those from a 3-methyl-3-buten-1-ol standard (Fig. 2C). *In vitro* activity measurement was attempted to determine the kinetic parameters of PMD_{sc} for MVA, but the enzyme activity was too low to determine the kinetic parameters (data not shown).

Structural analysis of a homologous PMD from *Staphylococcus epidermis* (PMD_{se}) (Barta et al., 2012) suggested that the diphosphate group is important for substrate binding even though it is not directly involved in the catalytic decarboxylation reaction



Fig. 3. Hydrolysis of IP and IPP by purified NudB or *E. coli* cell lysates. (A) IP hydrolysis. IP was hydrolyzed to isopentenol by *E. coli* cell lysates while isopentenol was not detected (*) from other two reactions with or without purified NudB. (B) Profile of IP and IPP concentrations in *in vitro* hydrolysis reactions of IPP by purified NudB.

(Supplementary Fig. S1). The importance of the diphosphate group in PMD_{se} activity implies that the monophosphorylated substrate (i.e. MVAP) might be better suited for decarboxylation than the substrate without any phosphate group (i.e. MVA).

3.3. Engineering of IPP-bypass pathway II and pathway optimization in E. coli

To verify the improved activity of PMD_{sc} for phosphorylated substrates (MVAP), an *in vitro* assay was performed with both MVA and MVAP. While isopentenol was not detected in the *in vitro* reaction, a detectable amount of IP was produced when MVAP was used as a substrate for PMD_{sc} . This result indicates that PMD_{sc} has higher decarboxylase activity towards MVAP than MVA and suggests that the phosphate group of MVAP does indeed enhance substrate binding and catalysis (Supplementary Fig. S1). The k_{cat} (0.14 s⁻¹) and K_m (0.99 mM) of PMD_{sc} toward MVAP (Supplementary Fig. S2) were about 35-fold lower and 8-fold higher than the reported k_{cat} (4.9 s⁻¹) and K_m (123 μ M) toward the native substrate (MVAPP), respectively (Krepkiy and Miziorko, 2004).

With a confirmation of promiscuous PMD_{sc} activity for MVAP, we constructed a new IPP-bypass pathway (pathway II in Fig. 1) by expressing AtoB, HMGS, HMGR, MK, and PMD_{sc} in *E. coli* (strain ARK2a, Table 1). Strain ARK2a produced 474.7 mg/L of isopentenol, a 558-fold improvement over the strain with pathway I (strain ARK3a). This new strain (strain ARK2a) achieved about 62.4% of the titer of the original isopentenol pathway (pathway O with strain ARK1a). It is noteworthy that IPP-bypass pathway II could produce isopentenol even without over-expressing any additional phosphatase that would hydrolyze the phosphate group in IP, which will be discussed in detail in the next section.

3.4. Identification of endogenous phosphatase for IP

The successful production of isopentenol via IPP-bypass pathway II suggested that endogenous E. coli phosphatases are capable of hydrolyzing IP to isopentenol. Initially, we hypothesized that IP might be hydrolyzed by promiscuous activities of Nudix hydrolases such as NudB in E. coli or an E. coli homolog of B. subtilis NudF, both of which were previously used to convert IPP to isopentenol (Chou and Keasling, 2012; Withers et al., 2007). In the original IPPdependent isopentenol pathway, the expression of NudB or NudF was essential for isopentenol production from IPP, and in vitro kinetic experiments of NudB showed that IPP was hydrolyzed by the enzyme (Chou and Keasling, 2012). However, the previous assay was based on the detection of the monophosphate formation without analyzing the final product by LCMS or GCMS, and it was not determined whether NudB hydrolyzes IPP by two consecutive hydrolysis reactions of two monophosphates or by a single hydrolysis reaction of a diphosphate group. We hypothesized that if NudB hydrolyzes IPP via the former fashion (i.e. two consecutive hydrolyses), both IP (intermediate) and isopentenol would be detected from an in vitro assay containing purified NudB and IPP. Interestingly, an in vitro assay of purified NudB with IPP produced only IP-no isopentenol was detected even after an extended incubation of 16 hours (Fig. 3A). Similarly, purified NudB could hydrolyze DMAPP to DMAP, but the final hydrolysis product, 3-methyl-2-buten-1-ol, was not detected (Supplementary Fig. S3A). In addition to NudB, NudF of B. subtilis, which was identified as a IPP hydrolase in a previous study (Withers et al., 2007), was also found to hydrolyze IPP to IP, but not to isopentenol (Supplementary Fig. S3B). On the other hand, it was confirmed in vitro that crude cell lysates of *E. coli* did hydrolyze IP to isopentenol (Fig. 3B). This result suggests that in the original isopentenol pathway. NudB hydrolyzed IPP to IP, but the following hydrolysis of IP to isopentenol was catalyzed by unknown endogenous phosphatase (s) in E. coli.

To identify the unknown endogenous phosphatase(s), phosphatase single gene knockout mutants were tested for their capability to hydrolyze IP to isopentenol. We reasoned IP hydrolysis to isopentenol would significantly decrease if the responsible IPhydrolyzing enzyme was absent in the knockout mutant strain. A total of 36 monophosphatase single gene knockout mutant strains were obtained from the Keio collection (Baba et al., 2006), and cell lysate from each individual strain was incubated with IP in vitro. Cell lysates from three single gene knockout mutant strains (Δagp , $\Delta aphA$ and $\Delta yqaB$) produced significantly less isopentenol than the average level of isopentenol produced by all strains tested including the wild type (Fig. 4A). Even after 26 h of incubation, the relative isopentenol level produced from cell lysates of Δagp , Δ *aphA* and Δ *yqaB* mutants were only 62%, 64% and 82% of the level from the wild type, respectively (Fig. 4B). It is noteworthy that cell lysates of all 36 mutants has some IP-hydrolyzing activity, suggesting that multiple endogenous phosphatases capable of hydrolyzing IP.

IP-hydrolysis efficiency significantly increased when one of these three genes, *aphA*, was overexpressed both in wild type and in the *aphA*-knockout mutant. In these strains, IP was completely converted into isopentenol immediately after addition of the IP to hydrolysates reactions (Fig. 4C). On the other hand, overexpression of the other two genes (*agp* and *yqaB*) showed relatively much slower IP hydrolysis rates (Fig. 4C), suggesting that *aphA* has much higher IP-hydrolysis activity than those of *agp* and *yqaB*. Co-expression of *aphA* along with pathway II (AtoB, HMGS, HMGR, MK and PMD_{sc}; strain ARK2aa) resulted in an isopentenol titer of 705 mg/L after 31 hours of incubation, which is about 20% higher than that of the strain without *aphA* overexpression (strain ARK2a; Fig. 5) and 83% of the maximum titer of the original isopentenol pathway (pathway O in



Fig. 4. Identification of endogenous phosphatases for IP. (A) Isopentenol concentration (μ M) in the cell lysates of monophosphatase mutants. A total of 36 mutants (diamonds)—including Δagp (solid circle), $\Delta yqaB$ (solid triangle), $\Delta aphA$ (solid square) mutants and wild type BW25113 (open circle)—were screened. The gray line represents the average (251.9 μ M) of isopentenol concentrations detected from all mutants. (B) Isopentenol converted from 1 mM IP by cell lysates of wild type (BW25113, solid circle), Δagp (open circle), $\Delta yqaB$ (open triangle) and $\Delta aphA$ (open square) mutants. (C) Isopentenol converted from 500 μ M IP by cell lysates of wild type (solid lines) or each mutant (dotted lines) with overexpression of the corresponding gene: agp (circle), yqaB (triangle) and aphA (square).



Fig. 5. Effect of *aphA* expression on isopentenol production in pathway II. Isopentenol from pathway II with (dark gray bar) or without *aphA* expression (white bar). Optical density of cell cultures at 600 nm (OD₆₀₀) for pathway II with (solid circle) or without *aphA* expression (open circle).

Fig. 1; strain ARK1a in Table 1; 836.9 mg/L). Although AphA is a membrane-bound protein whose overexpression frequently is detrimental and inhibits growth (Wagner et al., 2008), there was no significant growth difference between strains with or without *aphA*-overexpression (strains ARK2aa and ARK2a, respectively). Achieving a significant improvement in conversion of IP to isopentenol by *aphA* overexpression, we reasoned that the overall flux to isopentenol in strain ARK2aa could be further improved by increasing the activity of

 $\mathsf{PMD}_{\mathsf{sc}}$ toward MVAP. We thus focused on improving the promiscuous activity of $\mathsf{PMD}_{\mathsf{sc}}$ toward non-native substrates.

3.5. PMD engineering for improved activity toward mevalonate monophosphate

To engineer the active site of PMD_{sc} for the non-native substrate MVAP, we first identified amino acid residues in PMD putatively responsible for binding the native substrate (MVAPP). Since the only X-ray crystal structure of PMD_{sc} was solved without a bound substrate (Bonanno et al., 2001), the coordinates of MVAPP in the active site of PMD_{sc} were predicted by aligning the crystal structure of PMD_{sc} (PDB#: 1FI4) to that of the homologous PMD enzyme from S. epidermis (PMDse, PDB#: 4DPT) (Supplementary Fig. S4A). Crystal structures of PMD_{se} were solved with two substrate analogs: adenosine 5-[γ -thio]triphosphate (ATP γ S) and 6-fluoromevalonate 5-diphosphate (FMVAPP) (Barta et al., 2012). Alignment of PMD_{sc} and PMDse amino acid sequences showed 50% similarity by BLAST search and revealed conserved residues for catalysis and substrate binding (Supplementary Fig. S4B). However, in vivo isopentenol production with IPP-bypass pathway II and PMD_{se} (strain ARK4) was significantly reduced relative to PMDsc (11.3 mg/L vs 474.7 mg/L after 24 h), suggesting that the activity of PMDse toward MVAP could be much lower than that of PMD_{sc} . The k_{cat}/K_m ratios of PMD_{sc} and PMD_{se} are $4.0 \times 10^4 \, \text{s}^{-1} \, \text{M}^{-1}$ (Krepkiy and Miziorko, 2004) and $6.5 \times 10^5 \text{ s}^{-1} \text{ M}^{-1}$ (Barta et al., 2012), respectively, which indicates higher substrate specificity of PMDse for MVAPP. Increased substrate specificity in PMD_{se} could be attributed to the positively charged arginine at residue 193 (R193) (Barta et al., 2012). R193 of PMD_{se} is located within hydrogen bonding distance of the β -phosphate moiety of MVAPP and stabilizes the binding of MVAPP to the enzyme. On the other hand, PMD_{sc} has a neutral threonine residue in the homologous position (T209) instead of the positively charged arginine (Supplementary Fig. S4B), and this perhaps allows the promiscuity of PMD_{sc} towards the less negatively charged MVAP.

After we engineered the bypass pathway II with PMD_{sc}, an archaeal MVAP-specific decarboxylase was identified in Haloferax volcanii (PMD_{hv}) with much better kinetics for MVAP (K_m of 0.159 mM and k_{cat} of 3.5 s⁻¹ for MVAP; and no activity toward MVAPP) (Vannice et al., 2014). Unlike the conventional MVA pathway that supplies IPP via decarboxylation of MVAPP, the archaeal MVA pathway produces IPP via phosphorylation of IP, which is produced by decarboxylation reaction of MVAP similar to our bypass pathway II. Therefore, PMD_{hy} was expected to be a natural decarboxylase that can convert MVAP to IP in the IPPbypass pathway II. Surprisingly, however, no isopentenol production was detected when four pathway genes in the bypass pathway II (AtoB, HMGS, HMGR and MK) were expressed in vivo along with PMD_{hv} (strain ARK5; data not shown). An ATP-NADH coupled assay was also performed in vitro to detect the activity of PMD_{hv} toward MVAP, but no ATP hydrolysis activity was observed either. In the previous work where PMD_{hv} kinetics were determined, PMD_{hv} was overexpressed in its native host, *H. volcanii*, at 42 °C in salt-rich Hv-YPC media (containing 144 g of NaCl, 21 g of $MgSO_4 \cdot 7H_2O$, 18 g of $MgCl_2 \cdot 6H_2O$ and 4.2 g of KCl) (Vannice et al., 2014). Given that optimal growth temperatures and salt concentration in media of H. volcanii are different from those for E. coli, PMD_{hv} could have been expressed but inactive in E. coli. Nonetheless, it was noteworthy that four residues from PMD_{sc} that interact with β -phosphates of MVAPP were missing in PMD_{hv} between threonine 186 (T186) and glutamate 187 (E187) (Vannice et al., 2014). In other homologous PMD sequences from species with conventional MVA pathways, these missing residues are rich in serine and arginine, which facilitates interaction with the phosphoryl moieties of MVAPP and ATP. Therefore, analysis of residues near β -phosphate of MVAPP in three PMDs suggested that the activity of PMD_{sc} toward MVAP could be improved by redesigning the local electrostatic environment around the β phosphate of MVAPP.

Based on structural analysis of these three PMDs (PMD_{sc}, PMD_{se} and PMD_{hv}), four residues (K22, S155, S208 and T209) of PMD_{sc} adjacent to the β -phosphate of the MVAPP were selected for engineering (Supplementary Fig. S5). While the original substrate MVAPP has a net charge of -4, two alternative substrates. MVAP and MVA, have a net charge of -2 and 0, respectively. To compensate for this reduced negative charge, two serine residues (S155 and S208) were mutated to negatively charged glutamate (E), and the other two residues near the phosphate moiety (K22 and T209) were mutated to neutral methionine (M) and negatively charged aspartate (D), respectively. In addition, we constructed two more mutants, R74H and I145F (Fig. 6A), which were previously shown to increase activity of PMD_{sc} in the similar decarboxylation reaction for 3-hydroxy-3-methylbutyrate (3-HMB) to produce isobutene (Gogerty and Bobik, 2010). In vitro assay reactions using cell lysates of two serine-to-glutamate mutations (S155E and S208E) did not produce detectible amount of product, which suggests that these two mutations significantly reduced the activity of PMD_{sc} toward MVAP unlike the other mutants (data not shown). The K22M mutation increased $K_{\rm m}$ and decreased $k_{\rm cat}$, but



Fig. 6. Effect of mutations on isopentenol production with IPP-bypass pathway II. (A) location of R74 and I145 in PMD_{sc}. Blue meshes are essential residues for catalysis and substrate binding, and pink meshes are residues selected for mutagenesis. Electrostatic interactions were not clearly found between substrates and these residues and the distance between phosphate group of the substrate analog (6-fluoromevalonate 5-diphosphate (FMVAPP)) and R74 or 1145 residue were 12.5 Å or 15.0 Å, respectively. (B) Curve fittings and kinetics of PMD_{sc} wild type and four mutants (K22M, R74H, 1145F and T209D). (C) Isopentenol production from strains with pathway II containing different PMD mutants including wild type (WT, black square), R74H (open circle), 1145F (gray circle) and R74H/1145F double mutants (black circle).

Table 2

Kinetic parameters of PMD wild type, PMD_{sc} mutants, PMD_{se} , PMD_{hv} from other literatures.							
Name		$K_{\rm m}({ m mM})$	$k_{\rm cat}~({\rm s}^{-1})$	$k_{\rm cat}/K_{\rm m}~({\rm s}^{-1}~{\rm M}^{-1})$	% of WT	Substr	
PMD _{sc}	WT	0.99	0.14	1.4×10^2	100%		

Name		$K_{\rm m}$ (mM)	$k_{\text{cat}} (\mathrm{s}^{-1})$	$k_{\rm cat}/K_{\rm m}~({\rm s}^{-1}~{\rm M}^{-1})$	% of WT	Substrate	Reference
PMD _{sc}	WT	0.99	0.14	$1.4 imes 10^2$	100%		
	R74H	0.77	0.23	$3.0 imes 10^2$	220%		
	K22M	2.47	0.09	$3.5 imes 10^1$	25%	MVAP	This study
	T209D	0.99	0.13	1.3×10^{2}	98%		
	I145F	1.36	0.28	$2.0 imes 10^2$	147%		
PMD _{hv}		0.159	3.5	$2.2 imes 10^4$		MVAP	(Vannice et al., 2014)
PMD _{se}		0.009	5.9	$6.5 imes 10^{5}$		MVAPP	(Barta et al., 2012)
PMD _{sc}		0.123	5.4	$4.0 imes 10^4$		MVAPP	(Krepkiy and Miziorko, 2004)

the kinetic parameters of the T209D mutant were similar to those of the wild type (Table 2, Fig. 6B). Interestingly, the specificity of PMD_{sc} toward MVAP (k_{cat}/K_m) with R74H or I145F mutation was 220% and 147% of that of wild type, respectively. Although R74 and I145 are located near the active site, it is unlikely that these residues interact directly with substrates: distances from the α phosphate group of MVAPP are 12.5 Å and 15.0 Å, respectively (Fig. 6A). Therefore, the improved activity of the R74H and I145F mutants toward MVAP and 3-HMB suggests that these two mutations changed the conformation of the active site to accommodate less negatively charged substrates. Although R74H and I145F increased activity for MVAP and 3-HMB, two mutants did not show detectible hydrolysis activity on MVA.

After identifying two mutations in PMD_{sc} that improve activity toward MVAP, we prepared E. coli strains overexpressing four enzymes (AtoB, HMGS, HMGR, MK) along with one of three different PMD mutants including R74H (stain ARK2a_{M1}), I145F (strain ARK2a_{M2}), or the double mutant (strain ARK2a_{M3}) to see whether improved specificity for MVAP would increase isopentenol production in IPP-bypass pathway II. As shown in Fig. 6C, R74H (strain ARK2a_{M1}) resulted in significantly improved productivity (20.4 mg/L/hr) over wild type (15.9 mg/L/hr) through 30 hours of batch fermentation. The I145F mutation (strain ARK2a_{M2}), however, reduced isopentenol titer and productivity in vivo even though this mutation improved in vitro enzyme activity (Table 2). Interestingly, when these two mutations were combined (strain ARK 2a_{M3}), the titer and productivity were recovered to the comparable level to those of R74H, which suggests that R74H mutation was dominant over the I145F mutation.

Successful identification of PMD mutants that improve or significantly reduce isopentenol titer and productivity supports the hypothesis that the promiscuous activity of PMD toward MVAP is the current bottleneck of the IPP-bypass pathway II. Given the huge engineering space to explore various mutations that can potentially improve the activity of PMD toward MVAP, this result provides a clear opportunity to improve IPP-bypass pathway II for isopentenol production.

3.6. Effect of MVA levels on isopentenol production in the IPP-bypass pathway II

We successfully engineered IPP-bypass MVA pathways for isopentenol production and showed that pathway II could be improved by facilitating two limiting reactions: hydrolysis of IP and decarboxylation of MVAP to IP. Next, we targeted the "top" portion of the MVA pathway with engineering that would modulate pathway flux to MVA and tested how this variation affects isopentenol production in IPP-bypass pathway II. Previously, heterologous MVA pathways were constructed and tested with various combinations of HMGS and HMGR, and different pairs of HMGS-HMGR resulted in different levels of MVA and final isoprenoid titers (George et al., 2014; Pfleger et al., 2006; Pitera et al.,



Fig. 7. Effect of different "top" portions on isopentenol production in E. coli with pathway O or with pathway II. Four different "top" portions have different HMGS and HMGR sequences, which are original sequences from. S. cerevisiae (MevTo), codon-optimized sequences of S. cerevisiae (MevTco), sequences from S. aureus (MTSA) and sequences from *D. acidovorans* (MTDA). Isopentenol production was measured at 24 hours (white dotted bar) and at 48 h (gray bars).

2007; Ma et al., 2011). The MVA level was reported to affect MK activity by substrate inhibition (Ma et al., 2011), and therefore, optimizing MVA flux has been one approach to improve titers of isoprenoid products.

To evaluate the effects of MVA concentration in IPP-bypass pathway II, we reconstructed the original and the modified pathways with four different pairs of HMGS and HMGR in the "top" portion of the pathway (Fig. 1): non-codon optimized original sequences from S. cerevisiae genes (MevTo), E. coli-codon optimized sequences of S. cerevisiae genes (MevTco), HMGS and HMGR of Staphylococcus aureus (MTSA), and those of Delftia acidovorans (MTDA).

In accordance with the previous reports, the original IPPdependent isopentenol pathways showed different isopentenol titers depending on which pairs of HMGS and HMGR were used (Fig. 7). Analysis of intracellular metabolites confirmed that expression of different pairs of HMGS and HMGR indeed resulted in various intracellular MVA concentrations in strains with both pathway O and pathway II (Supplementary Fig. S6A). Intriguingly, isopentenol titers from the strains containing IPP-bypass pathway II did not change much when the pairs of HMGR and HMGS are changed (Fig. 7), and similar levels of IP were also observed in the strains with pathway II (Supplementary Fig. S6D). This "insensitivity" of the isopentenol titer to various "top" portions and a similar level of IP in pathway II strains suggests that the determining factor of isopentenol production in pathway II could be the PMD activity toward MVAP rather than upstream pathway efficiency.

In addition, metabolite analysis showed that strains with pathway O or pathway II accumulated significantly high levels of IPP or MVAP, respectively, regardless of intracellular MVA concentrations (Supplementary Fig. S6). Interestingly, MVAP was accumulated to considerably higher concentrations than that of IPP (100~200 mM for MVAP vs 30-60 mM for IPP) without any significant toxicity, which is consistent with the previous report that MVAP is not inhibitory to cell growth (Martin et al., 2003).

3.7. Relief of IPP-toxicity in the bypass pathway II

Previous studies showed that the performance of the original MVA pathway was sensitive to MK expression levels: low MK expression resulted in attenuated flux to IPP and isopentenol, but high levels led IPP accumulation and resulted in growth inhibition (George et al., 2014, 2015). Interestingly, growth was restored when NudB was overexpressed in IPP-accumulating strain to relieve IPP-toxicity. In the current study, we demonstrated that NudB could hydrolyze IPP to IP, but not further to isopentenol (Fig. 3), suggesting that IPP has more detrimental effects on growth than does IP.

Since the bypass pathway II does not produce IPP, we hypothesized that the pathway would be insensitive to changes in MK expression and free from related toxicity. To compare growth and isopentenol production in the original and IPP-bypass pathway (pathway O and pathway II) under IPP- or IP-accumulating conditions, respectively, two modifications were made to the strains ARK1a and ARK2a (Supplementary Fig. S7). First, to achieve a moderate level of MK expression in the control strains, we removed the promoter previous added for MK overexpression in the medium copy plasmids JBEI-12056 and JBEI-9310. With this engineering, MK was expressed at a moderate level as the forth enzyme in the operon containing three enzymes for the top portion of the MVA pathway, and it resulted strains ARK1e (harboring JBEI-6818 and JBEI-6833) and ARK2e (harboring JBEI-12051 and [BEI-9314). Second, to achieve very high MK expression level, an additional copy of MK was added to the high copy plasmids, JBEI-6833 and JBEI-9314, resulting ARK1f and ARK2f, respectively (Supplementary Fig. S7). Confirming the previous results (George et al., 2014), balancing flux in the upstream pathway was critical for growth and isopentenol production (Fig. 8). Growth and isopentenol production of ARK1f was significantly reduced showing sensitivity to expression levels of MK, but strains with pathway II was insensitive to downregulation (ARK2f) or up-regulation (ARK2e) of MK. and free from burden of IPP accumulation (Fig. 8).

3.8. The effect of limited aeration on isopentenol production via IPPbypass pathway

After characterizing the IPP-bypass pathway II in E. coli (strain ARK2a), we tested whether this pathway would have any advantage over the original pathway under ATP-limited conditions. In general, ATP is most efficiently supplied via oxidative phosphorylation with oxygen as a final electron acceptor. As a result, aeration has been an important operation in industrial-scale fermentation, especially when ATP-demanding isoprenoid biosynthetic pathways are exploited. However, the aeration cost is usually one of the largest portions (up to 26%) of the overall utility cost, and the cost would be on the order of \$60 million per year in a plant that processes 2000 MT of dry biomass per day (Clark and Blanch, 1997). Moreover, oxygen mass transfer is limited in large-scale fermenters, and this potentially creates a local micro-aerobic or anaerobic environment during fermentation. Therefore, the development of fermentation processes with reduced aeration rates can significantly reduce production cost and improve process efficiency. With this goal in mind, we investigated the impact of reduced aeration on isopentenol production with pathways O and II, which require 3 ATPs and 2 ATPs, respectively, to produce one molecule of isopentenol. To provide different aeration rates, we prepared a 50-mL cell culture with an OD₆₀₀ of 0.6–0.7, split into two 25-mL cell cultures in 250-mL flasks, and continued to incubate at 30 °C for induction (0.5 mM IPTG) at two different shaking speeds (30 rpm and 200 rpm).

Fig. 9A shows that the isopentenol titer of pathway O (strain ARK1a) was more significantly affected when aeration was limited by lowering the shaking speed from 200 rpm to 30 rpm. With a reduced aeration, strain ARK1a produced only 22% of the initial



Fig. 8. IPP toxicity in Pathway O. (A) Growth of four strains containing Pathway O without (black, square) or with expression of additional MK (white square); Pathway II without (black, circle) or with expression of additional MK (white circle). (B) Isopentenol production from four strains containing Pathway O without or with expression of additional MK; Pathway II without or with expression of additional MK.

titer at 200 rpm after 16 h-fermentation (Fig. 9B). The bypass pathway II (strain ARK2a), however, produced 40% at 16 h and up to 60% of the titers under the higher aeration conditions at 24 h. It is noteworthy that the OD₆₀₀ of strain ARK1a was higher than that of strain ARK2a under poor aeration condition (at 30 rpm). A better growth but significantly less isopentenol production of strain ARK1a suggests that the heterologous MVA pathway may compete for ATP with other essential cellular processes related to the growth, and when ATP supply is limited (i.e. under poor aeration conditions), strain ARK1a might reduce the carbon flux to the MVA pathway to reduce the energy usage for this ATPconsuming heterologous pathway. The strain with pathway II (strain ARK2a), however, produced a similar or even higher level of isopentenol under limited aeration conditions after 16 h or 24 h of fermentation, respectively (Fig. 9A and Supplementary Fig. S8). This result also suggests that the bypass pathway II would be more robust when aeration is limited, and a reduced ATP demand in strain ARK2a is possibly beneficial to the strain under oxygenlimited conditions. Therefore, more economic production of isopentenol could be feasible via the ATP-saving IPP-bypass pathway II by reducing aeration costs for large scale fermentation.

4. Conclusion

Isopentenol is a potential gasoline alternative and a precursor of commodity chemicals such as isoprene. In this study, we reported our efforts to remove "IPP-dependency" of the original MVA pathway and to overcome limitations intrinsic to IPP accumulation and "unnecessary" consumption of ATPs for isopentenol production. By implementing two previously unidentified activities of PMD_{sc} and AphA, we demonstrated that considerable



Fig. 9. Effect of reduced aeration conditions on isopentenol production in *E. coli*. (A) Isopentenol titers (per OD₆₀₀) of two strains with pathway O or with pathway II under higher (200 rpm) or lower (30 rpm) aeration conditions. (B) Relative isopentenol titers (total) of two pathways under lower aeration conditions (30 rpm) compared to those under higher aeration conditions (200 rpm).

isopentenol titers could be achieved without producing IPP via the pathway II.

The IPP-bypass pathway II was shown to be a robust alternative to the original pathway (pathway O) for isopentenol production. This modified pathway was insensitive to both MVA level and MK expression level, and reduced the engineering burden to balance the upstream MVA pathway and IPP toxicity. Most significantly, the IPP-bypass pathway II was more competitive when aeration was limited, which would significantly lower operational costs for aeration in a large scale fermentation.

Finally, in this report, we found that the promiscuous activity PMD is rate-limiting. The identification of PMD as the rate-limiting step in these bypass pathways provides clear engineering opportunities. Although we constructed a few PMD mutants with improved activity toward MVAP, more concerted efforts to engineer PMD promiscuity or identify homologous enzymes should yield additional increases in isopentenol yield and productivity. With further engineering, these bypass pathways will provide valuable platforms for the energetically-favored production of isopentenol, isoprene, and related C_5 compounds.

Acknowledgments

We thank Dr. Miziorko at University of Missouri-Kansas City for providing a plasmid containing the PMD_{se} gene and Dr. Konda at JBEI for discussion on techno-economic impact of this work. This work was part of the DOE Joint BioEnergy Institute (http://www. jbei.org) supported by the U.S. Department of Energy, Office of Science, Office of Biological and Environmental Research, through contract DE-AC02-05CH11231 between Lawrence Berkeley National Laboratory and the U.S. Department of Energy. The United States Government retains and the publisher, by accepting the article for publication, acknowledges that the United States Government retains a non-exclusive, paid-up, irrevocable, world-wide license to publish or reproduce the published form of this manuscript, or allow others to do so, for United States Government purposes.

Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.ymben.2015.12.002.

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