Designing and Constructing a Novel Artificial Pathway for Malonic Acid Production Biologically

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Malonic acid is used as a common component of many products and processes in the pharmaceutical and cosmetic industries. Here, we designed a novel artificial synthetic pathway of malonic acid, in which oxaloacetate, an intermediate of cytoplasmic reductive tricarboxylic acid (rTCA) pathway, is converted to malonic semialdehyde and then to malonic acid, sequentially catalyzed by α-keto decarboxylase and malonic semialdehyde dehydrogenase. After the systematic screening, we discovered the enzyme oxaloacetate decarboxylase Mdc, catalyzing the first step of the artificially designed pathway in vitro. Then, this synthetic pathway was functionally constructed in cellulolytic thermophilic fungus Myceliophthora thermophila. After enhancement of glucose uptake, the titer of malonic acid achieved 42.5 mg/L. This study presents a novel biological pathway for producing malonic acid from renewable resources in the future.

Keywords: malonic acid, α-keto decarboxylase, oxaloacetic acid, metabolic engineering, Myceliophthora thermophila

INTRODUCTION

Malonic acid, formally known as propanedioic acid, is widely used in manufacturing processes, such as the petrochemical, pharmaceutical, and cosmetic industries. Traditionally, malonic acid is produced from fossil resources via petrochemical processes. The byproduct sodium cyanide is dangerous and unfriendly to the environment (Hildbrand and Pollak, 2012; Werpy et al., 2004; Klikar et al., 2017). With increasing concerns on energy and environmental problems, the production of malonic acid by microbial fermentation via bioconversion of renewable feedstock has generated considerable interest worldwide. However, due to a lack of knowledge about the malonic acid synthesis pathway, there is little progress on the biological production of malonic acid.

Recent advances in synthetic biology and computational biology have enabled designing novel and specific metabolic pathways for desired chemicals (Atsumi et al., 2008). However, importing and overexpressing non-native pathways in microbes may lead to metabolic imbalance, resulting in none or low titer of target products. It is critical to design artificial synthetic pathways that are compatible with the host. At present, two non-natural metabolic pathways have been sought and introduced into microbes, using malonic semialdehyde (MSA) or malonyl-CoA as the precursor (Table 1) (Song et al., 2016; Dietrich et al., 2017; Chae et al., 2020). The first one uses MSA from the deamination reaction of β-alanine as the precursor. In Escherichia coli, β-alanine pyruvate transaminase from Pseudomonas aeruginosa was overexpressed to convert β-alanine to produce MSA. Then, succinate semialdehyde dehydrogenase encoded by E. coli yndE was used to catalyze the reaction of MSA to malic acid, and the titer of malonic acid reached 3.60 g/L by fed-batch fermentation (Song et al., 2016). Another pathway is dependent
TABLE 1 | Summary of microbial production of malonic acid from glucose.

| Microbe         | Precursor | Description | Titer   | Fermentation | Refs       |
|-----------------|-----------|-------------|---------|--------------|------------|
| Escherichia coli | β-Alanine | +ppc, aspA, and ynl from E. coli; +panD from Corynebacterium glutamicum; | 0.45 g/L | Fed batch    | Song et al. (2016) |
| E. coli         | Acyl-CoA  | +ehd3 mutant from Saccharomyces cerevisiae | 3.6 g/L | Shake flask  | Dietrich et al. (2017) |
| Pichia kudriavzevii | Acyl-CoA | +ehd3 mutant from S. cerevisiae; +Annmae1 from Aspergillus niger | 82.3 mg/L | Shake flask  | Dietrich et al. (2017) |
| Myceliophthora thermophila | Oxaloacetate | +mdc from Ogataea parapolymorpha; +ynl from E. coli; +glt-1 from Neurospora crassa | 76 mg/L | Shake flask  | This study |

Note. “+” represents overexpression of target gene; “Δ” represents disruption of target gene.

on the fatty acid synthesis pathway, using malonyl-CoA as the precursor. Matthew et al. reported that acyl-CoA hydrolase encoded by ehd3 from Saccharomyces cerevisiae can hydrolyze malonyl-CoA to malonic acid. During overexpression of EHD3 in S. cerevisiae, Pichia kudriavzevii, and E. coli, malonic acid in the cultures was detectable. After improvement of binding affinity of acyl-CoA hydrolase to malonyl-CoA by protein engineering, production of malonic acid was increased to 0.0823 g/L by engineered E. coli (Dietrich et al., 2017). Both current synthetic pathways of malonic acid require a number of enzymes to convert glucose to target acid and are complex, which might lead to a low titer of malonic acid. So more research is needed to improve the production of this dicarboxylic acid, including novel biosynthetic pathway design, key enzymatic engineering, and even alternative hosts.

Filamentous fungi are classical predominant microorganisms for organic acid production, such as citric acid, malic acid, and lactic acid (Dave and Punekar, 2015; Kirimura et al., 2016; Li et al., 2020a). The thermophilic filamentous fungus Myceliophthora thermophila (synonym: Thermothelomyces thermophila), possessing the ability to efficiently degrade plant biomass, represents a potential library of new industrial applications, including producing thermo-tolerant and efficient cellulolytic enzymes and synthesizing biochemicals and biofuels directly from plant biomass (Berka et al., 2011; Yang et al., 2015; Singh, 2016). M. thermophila has been employed to synthesize bio-based products, such as malic acid, fumaric acid, and bioethanol, through metabolic engineering using CRISPR/CAS9 system (Liu et al., 2017; Gu et al., 2018; Li et al., 2020a; Li et al., 2020b). In this study, we designed a novel synthetic pathway of malonic acid and screened the key enzyme, oxaloacetate decarboxylase, in vitro. In this pathway, oxaloacetic acid (OAA) can be converted to MSA and then to malonic acid and sequentially catalyzed by α-keto decarboxylase and MSA dehydrogenase. Moreover, this artificial synthetic pathway was functionally constructed in M. thermophila to produce malonic acid from glucose.

MATERIALS AND METHODS

Strains and Culture Conditions

M. thermophila ATCC 42464 was obtained from the American Type Culture Collection (ATCC). The wild-type strain and its mutants were grown on Vogel’s minimal medium supplemented with 2% glucose (MM medium) at 35°C for approximately 12 days to obtain mature conidia, and antibiotics were added when needed for transformant screening.

E. coli DH5α and E. coli BL21 (DE3) were cultivated in Luria–Bertani medium supplemented with 100 μg/ml of ampicillin for plasmid selection.

Vector Construction

For overexpressing target genes in E. coli, the codon-optimized genes encoding oxaloacetate decarboxylase and MSA dehydrogenase were synthesized by GENEWIZ (Suzhou, China) and inserted between NdeI and XhoI of the plasmid pET-21a (+).

For the construction of plasmids overexpressing target genes in M. thermophila, codon-optimized mdc (XP_013934857.1) from Ogataea parapolymorpha, under control of the strong constitutive promoter of Pap amplified from the plasmid pPK2, was inserted between the BglII and BamHI sites of pAN52-PtrpC-Bar-PMtgpdA to generate the vector pAN52-Pap-mdc, using the NEB Gibson Assembly Kit. Similarly, the PCR fragments of the strong constitutive promoter of pgk (MYCTH_2316240) from M. thermophila and the opening reading fragment of ynl from E. coli were inserted between the BglII and BamHI sites of pAN52-PtrpC-bar-PMtgpdA to generate the plasmid pAN52-PtrpC-bar-PMTgpdk-ynl. Under control of the strong constitutive promoter of pgk (MYCTH_2316240) of M. thermophila, Anmae1 encoding the C4-dicarboxylate transporter from Aspergillus niger were inserted between the BglII and BamHI sites of pAN52-PtrpC-neo-PMtgpk to form the vector pAN52-PtrpC-neo-PMtgpgk-Anmae1. The PCR fragments of glucose transporter-encoding genes glt-1 from Neurospora crassa were inserted between the Spel and BamHI sites of pAN52-PtrpC-bar-PMtgpdA to generate the plasmid pAN52-PtrpC-bar-PMtgpdA-Anmae1.

All vectors were constructed using E. coli DH5α. The target genes cloned into shuttle vectors were sequenced to verify the authenticity of the plasmid construction. All the primers used for plasmid construction are listed in Supplementary Table S1.

Recombinant Proteins Expression and Purification

E. coli BL21 (DE3) strains harboring expression plasmids were grown at 37°C in 100 ml of LB media supplemented with 100 μg/
ml of ampicillin to an optical density OD<sub>600</sub> of 0.4–0.5. Then, 0.4 mM of isopropyl-β-D-thiogalactopyranoside (IPTG) was supplemented, and the cells were cultivated at 16°C for 19 h. Cells were harvested by centrifugation at 4°C, resuspended in 50 mM of Tris-HCl (pH 7.5) buffer, and crushed with an ultrasonus crusher. After centrifugation, the supernatant was collected and loaded onto a Ni<sup>2+</sup>-NTA-agarose column pre-equilibrated with binding buffer (50 mM of TAE (Tris-acetate-EDTA) buffer, 300 mM of NaCl, and 20 mM of imidazole, pH 7.5). The retained proteins were recovered with elution buffer (20 mM of Na<sub>2</sub>HPO<sub>4</sub>–NaH<sub>2</sub>PO<sub>4</sub>, pH 7.4, 500 mM of NaCl, and 500 mM of imidazole). In order to eliminate salt and imidazole, the eluted fraction was filtered by an ultrafiltration centrifugal tube. The purified protein was stored at –20°C. The purity of the enzymes was analyzed by sodium dodeyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE), and protein concentration in supernatants was measured using a Bio-Rad protein assay kit based on absorbance at 590 nm, using bovine serum albumin as the standard.

In Vitro Enzyme Reaction

To synthesize malonic acid from OAA in vitro, the reaction mixture (2 ml) containing 50 mM (pH 7.5) of Tris–HCl, 1 mM of NAD<sup>+</sup>, 1 mM of MgSO<sub>4</sub>, 1 mM of thiamine pyrophosphate (TPP), excessive purified YneI (approximately 40 µl), and 100 µl of purified Mdc was used. The assay was started by the addition of 1 mM of OAA and was immediately monitored at 340 nm for 5 min. Subsequently, the reactants were analyzed by gas chromatography/mass spectrometry (GC/MS).

Gas Chromatography–Mass Spectrometry Assay

Samples (reaction products and malonic acid standard) were dried using a CentrVap vacuum concentrator. Then, the dried samples were dissolved in 20 µl of pyridine solution containing 20 mg/ml of methoxyamine hydrochloride and incubated for 90 min at 30°C. After that, 50 µl of the MSTFA reagent (containing 1% TMCS, v/v) was added to the sample aliquots, mixed well, and then incubated for 1 h at 37°C. The mixture was assayed by GC/MS as described previously (Mao et al., 2021).

Analytical Method of Sugar

Residual glucose in the culture was determined by high-performance liquid chromatography (HPLC) equipped with a Waters (Milford, MA, USA) 2414 refractive index detector and an Aminex HPX-87H column (Bio-Rad Laboratories, Hercules, CA, USA) at 35°C. H<sub>2</sub>SO<sub>4</sub> measuring 5 mM was employed as the neutralizing agent at a final concentration of 20 g/L to keep the pH at approximately 6.0.

RESULTS AND DISCUSSION

Design of an Artificial Biosynthetic Pathway of Malonic Acid

A novel artificial biosynthetic route was devised to produce malonic acid with OAA as the substrate. In this route, OAA is converted to malonic acid by a-keto decarboxylase (Kdc) and then to malonic acid (Figure 1A). Kdc is the crucial enzyme step required to divert flux from OAA to malonic acid, which is widespread in plants, yeasts, and fungi. Kdc has been widely used in the non-native synthetic pathway of biofuels, in which Kdc converted 2-keto acids from amino acid biosynthesis pathways to aldehyde and then catalyzed by dehydrogenase to produce alcohols, including 1-propanol, 1-butanol, and isobutanol (Atsumi et al., 2008). Substantial numbers of Kdc have been identified and characterized. Moreover, the protein engineering and method of high-throughput screening have been developed to improve the thermostability, substrate specificity, and activity of Kdc. Therefore, it was possible to find a Kdc to catalyze OAA to MSA. In addition, another key enzyme MSA dehydrogenase has been screened, and succinate semialdehyde dehydrogenase encoded by yneI from E. coli has been used for catalyzing MSA to malonic acid (Fuhrer et al., 2007; Sullivan and Holyoak, 2007; Zheng et al., 2013; Song et al., 2016). Therefore, the novel pathway for synthesizing malonic acid was theoretically feasible.

In microbes, OAA is the intermediate of the cytoplasmic reductive tricarboxylic acid (rTCA) pathway and mitochondrial TCA cycle. Assuming that glucose is channeled to OAA via the glycolysis pathway and TCA pathway in turn, 1 mol of glucose produces 2 mol of OAA and 2 mol of NADH, accompanied by fixation of 2 mol of CO<sub>2</sub> via carboxylation of pyruvate catalyzed by pyruvate carboxylase. The formed 2 mol of OAA can be converted to 2 mol of malonic acid and 2 mol of NADH. In total, 1 mol of glucose produces 2 mol of malonic acid and 4 mol of NADH, by which it achieves the theoretical maximum yield of 2 mol/mol of glucose (Figure 1B).
FIGURE 1 | The artificial synthetic pathway of malonic acid. (A) Oxaloacetic acid is converted to malonic semialdehyde by α-keto decarboxylase and then malonic acid by malonic semialdehyde dehydrogenase. (B) The theoretical maximum yield of malonic acid from glucose.

FIGURE 2 | The screening of Kdc using multi-enzyme system in vitro. (A) Summary of screening α-keto decarboxylase. (B) Sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) of purified Mdc (oxaloacetate decarboxylase) and Ynel (malonic semialdehyde dehydrogenase). Lane M, molecular weight marker; Lane 1, elution fraction containing purified Mdc (predicted molecular mass of the monomer, 62.9 kDa); and Lane 2, elution fraction containing purified Ynel (predicted molecular mass of the monomer, 50 kDa). (C) Time course of absorbance increase of NADH-coupled assay at 340 nm. Red: the reaction mixture only containing Ynel. Blue: the reaction mixture containing Mdc and Ynel. (D) Specific activity of purified single Mdc.
net NADH can be used for the synthesis of cell compounds and energy metabolites.

**Synthesis of Malonic Acid by Artificial Synthetic Route In Vitro**

TPP-dependent keto decarboxylase (Kdc) is a critical enzyme in malonic acid production, and some members have been used for the production of keto acid-derived alcohols (Iding et al., 1998). During the reaction of decarboxylation of OAA, β-carboxylate is easier to be replaced by hydrogen atom than α-carboxylate, which might result in difficulty of mining the highly active enzyme (Sullivan and Holyoak, 2007). To test the capability of using OAA as a substrate, five Kdc candidates, including Thi3, Aro10 from *S. cerevisiae* (Vuralhan et al., 2003), Pdc from *Clostridium acetobutylicum* (Iding et al., 1998), Kivd from *Lactococcus lactis* (Plaza et al., 2004), and Mdc from *O. parapolymorpha*, together with succinate semialdehyde dehydrogenase-encoding YneI from *E. coli* were heterologically overexpressed in *E. coli* BL21 (DE3) as an N-terminal His*6*-tagged protein. After induction of protein expression, the target proteins were purified and characterized. The NADH-coupled enzyme assay was chosen for the characterization of these purified Kdc enzymes (Figures 2A,B). As shown in Figure 2C, the only reaction mixture containing Mdc and excessive YneI exhibited increased absorbance at 340 nm. GC/MS analysis further revealed that purified Mdc and YneI catalyzed OAA to produce malonic acid (Figure 3), and the concentration of malonic acid was up to 8.1 mg/L. These data indicated that Mdc had the ability to catalyze OAA to MSA and the specific activity achieved 9.37 μmol/min/g (Figure 2D).

**The Artificial Malonate Biosynthetic Pathway Can Function in Myceliophthora thermophila**

Thermophilic fungus *M. thermophila* has been engineered to produce C4-dicarboxylic acids, malic acid, and fumaric acid, by cytoplasmic TCA pathway (Gu et al., 2018; Li et al., 2020a), which can provide precursor OAA for producing malonic acid. Therefore, in order to test whether this artificial malonic pathway can function in vivo, both *mdc* and *yneI* were integrated into the genome of *M. thermophila* wild-type strain, under the control of the strong constitutive promoters of *ap* (transcriptional enhancer factor) and *pgk*, respectively (Figure 4A). After confirmation via PCR analysis, the engineered strains were tested for the production of malonic acid with glucose as the carbon source. Malonic acid in the culture of the strain of SG214 was detected by GC/MS assay, and the titer achieved 40.8 mg/L (Figure 4B), indicating that the artificial malonate synthetic pathway we designed can work functionally in vivo.
presented a promising approach to produce malonic acid direct from renewable plant cell biomass in this cellulolytic thermophilic fungus in the future.

No Obvious Improvement was Obtained by Overexpression of Glucose Transporter and a Predicted Exporter of Malonic Acid

Rapid substrate supply was recognized as one strategy to maintain a high flux of reaction, which is a prerequisite for efficient cell factory production of biochemicals. Glucose transporter gene glt-1 from N. crassa has been systematically characterized and used for improving the production of commodity chemicals, such as malic acid and ethanol (Chen et al., 2017; Wang et al., 2017; Li et al., 2020b). Therefore, to improve the malonic acid production, gene glt-1 was incorporated into the M. thermophila SG214 strain, under the strong constitutive promoter of gpdA (MYCTH_2311855). After confirmation of the presence of the transgene by PCR analysis, the production of malonic acid by resultant strain SG317 was assayed. As shown in Figures 4B,C, the introduction of glt-1 enabled faster glucose utilization and malonate production, while titer of malonic acid (42.5 mg/L) exhibited a slight increase compared with the parental strain SG214 (Figure 4B). These data suggested that glucose absorbability is not the key bottleneck for the current version of the cell factory, and other strategies are needed to increase the production of malonic acid.

Previous research on metabolic engineering of chemical products showed that effective export of target products is the key factor of efficient production of desirable products, and overexpression of exporters leads to the improved synthesis of organic acids, such as malic and fumaric acid (Chen et al., 2017). It was reported previously that Anmae1, a C4-dicarboxylate transporter from A. niger, has the capability to export malonic acid, and it has been introduced into P. kudriavzevii for increased malonate production (Dietrich et al., 2017). Disappointingly, when the gene Anmae1 was overexpressed in the strain of SG214, under the control of the constitutive promoter of pgk (MYCTH_2316240), malonic acid production (41.3 mg/L) and glucose consumption rate by the resultant strain SG425 showed no significant increase, compared with the parent strain.

In order to improve the performance of cell factories for malonic acid production, more strategies of metabolic modification would be needed in the future, such as improvement of activity of key enzymes, disruption of branch points of the synthetic pathway, and enhancement of precursor pool size. In addition, intracellular high

\[ \text{FIGURE 4 | Introduction of malonic acid synthesis pathway in Myceliophthora thermophila. (A) Overall scheme of malonic acid synthesis in } M. \text{ thermophila. Time course of glucose consumption (B) and malonic acid production (C) by } M. \text{ thermophila mutants grown on glucose medium. WT, } M. \text{ thermophila wild-type strain; strain SG214 containing mdc and yneI; Strain SG317 containing mdc, yneI, and glt-1; Strain SG425 containing mdc, yneI, and Anmae1; PEP, phosphoenolpyruvate; GLUT, glucose transporter; PYC, pyruvate carboxylase; cMDH, cytoplasm malate dehydrogenase; Fum, fumarase; Fr, fumarate reductase; TCA, tricarboxylic acid cycle; PDH, pyruvate dehydrogenase; CS, citrate synthase; ACO, aconitase; IDH, isocitrate dehydrogenase; KGD, } \alpha \text{-ketoglutarate dehydrogenase complex; LSC, succinyl-CoA synthetase; SDH, succinate dehydrogenase; MDH, mitochondrial malate dehydrogenase.} \]
concentrations of malonate can competitively inhibit succinate dehydrogenase activity, which would lead to metabolic disturbance of host cells. Increasing host cell tolerance to malonate may be another strategy for the efficient production of malonic acid.

CONCLUSION

In this study, we designed a novel artificial pathway for malonic acid biosynthesis with OAA as the substrate. The pathway was first tested in vitro. OAA can be converted to MSA by α-keto decarboxylase (Mdc) and then to malonic acid by the second enzyme (Ynrl). Then, using a cellulolytic fungus M. thermophila as the host, this novel pathway was functionally constructed in vivo. The present study provides a new potential option to produce malonic acid from glucose and even plant biomass in the future.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding authors.

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AUTHOR CONTRIBUTIONS

CT and JL designed the experiments. SG, ZZ, YY, and JL performed the metabolic engineering experiments and analyze the data. CT, JL, and SG wrote the manuscript. All authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

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