Development of a promising microbial platform for the production of dicarboxylic acids from biorenewable resources

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Abstract

Background: As a sustainable industrial process, the production of dicarboxylic acids (DCAs), used as precursors of polyamides, polyesters, perfumes, plasticizers, lubricants, and adhesives, from vegetable oil has continuously garnered interest. Although the yeast Candida tropicalis has been used as a host for DCA production, additional strains are continually investigated to meet productivity thresholds and industrial needs. In this regard, the yeast Wickerhamiella sorbophila, a potential candidate strain, has been screened. However, the lack of genetic and physiological information for this uncommon strain is an obstacle that merits further research. To overcome this limitation, we attempted to develop a method to facilitate genetic recombination in this strain and produce high amounts of DCAs from methyl laurate using engineered W. sorbophila.

Results: In the current study, we first developed efficient genetic engineering tools for the industrial application of W. sorbophila. To increase homologous recombination (HR) efficiency during transformation, the cell cycle of the yeast was synchronized to the S/G2 phase using hydroxyurea. The HR efficiency at POX1 and POX2 loci increased from 56.3% and 41.7%, respectively, to 97.9% in both cases. The original HR efficiency at URA3 and ADE2 loci was nearly 0% during the early stationary and logarithmic phases of growth, and increased to 4.8% and 25.6%, respectively. We used the developed tools to construct W. sorbophila UHP4, in which β-oxidation was completely blocked. The strain produced 92.5 g/l of dodecanedioic acid (DDDA) from methyl laurate over 126 h in 5-l fed-batch fermentation, with a productivity of 0.83 g/l/h.

Conclusions: Wickerhamiella sorbophila UHP4 produced more DDDA methyl laurate than C. tropicalis. Hence, we demonstrated that W. sorbophila is a powerful microbial platform for vegetable oil-based DCA production. In addition, by using the developed genetic engineering tools, this emerging yeast could be used for the production of a variety of fatty acid derivatives, such as fatty alcohols, fatty aldehydes, and ω-hydroxy fatty acids.

Keywords: Wickerhamiella sorbophila, Alkane-assimilating yeast, Microbial platform, Dicarboxylic acid, Dodecanedioic acid, Vegetable oil, Genetic engineering tool

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Background

Industrial biotechnology, mainly known as white biotechnology in Europe, is a promising solution to replace the production of petroleum-based products that are widely used in daily life. Among various applications, the biotransformation of dicarboxylic acids (DCAs) from renewable resources has attracted much attention. DCAs are high value-added chemicals used as raw materials for a variety of products, such as polyamides (PAs), polyesters, perfumes, plasticizers, lubricants, and adhesives [1]. In particular, medium-chain DCAs (chain length 10–12), such as sebacic acid (C₁₀) and dodecanedioic acid (C₁₂), serve as precursors for PA 610 (nylon 6, 10) and PA 612 (nylon 6, 12) synthesis.

Alkane-assimilating yeasts are hosts for DCA biotransformation. Candida tropicalis with a strong ω-oxidizing ability is a representative of this group of yeasts. These microorganisms convert terminal methyl groups of alkanes and fatty acids into carboxylic groups via ω-oxidation and metabolize the corresponding fatty acids and DCAs via the β-oxidation pathway (Fig. 1). Several studies have focused on increasing DCA yield and productivity, for example, by blocking β-oxidation [2, 3] and enhancing ω-oxidation [4, 5] in C. tropicalis. This led to the industrialization of the DCA bioprocess using paraffin (alkane) [6]. However, considering the limited availability of petroleum-based resources, such as alkanes, sustainable production of DCA from vegetable oil-based resources is necessary. Among these, methyl laurate (C₁₂) is an attractive sustainable resource. It is obtained from the esterification of coconut and palm kernel oil—rich in lauric acid (C₁₂)—and is easily accessible for bioprocessing, since it is present in liquid form. Furthermore, medium-chain fatty acid methyl esters, including methyl laurate, are treated as by-products in the biodiesel industry, which relies on various vegetable oils to lower the cetane number associated with the ignition quality of fuel [7]. Consequently, DCA bioprocessing with vegetable oil-based feedstock as a starting material has been currently developed [8]. Unlike alkanes, however, biotransformation using vegetable oil-derived resources poses several difficulties. Cellular toxicity is one of the major obstacles that decrease the overall productivity of the bioprocess. In particular, medium-chain fatty acids (chain length 10–12) are highly toxic compounds, which cause cell membrane stress, disruption of the electron transport system, acidification of the cytoplasm, and production of reactive oxygen species [9, 10]. Medium-chain fatty acid methyl esters, such as methyl decanoate and methyl laurate, are hydrolyzed by extracellular lipases or esterases during the conversion phase and the resulting fatty acids cause cellular toxicity [11]. Because of its toxicity, various concentrations of DCA are produced, depending on the carbon chain length of fatty acids or that of methyl esters (Table 1).

We previously isolated a novel alkane-assimilating yeast Wickerhamiella sorbophila from wastewater of a petrochemical factory and attempted to use it to produce DDDA using methyl laurate [12]. The growth of W. sorbophila strain capable of ω-oxidation in the presence of mixed fatty acid methyl esters (C₁₀–C₁₆) was more stable than that of C. tropicalis and Yarrowia lipolytica [12]. Furthermore, it is a haploid and nonpathogenic yeast, with the associated advantages of genetic engineering and industrial safety [12]. Recently, whole-genome sequencing of W. sorbophila was completed, allowing easy access to the genetic information (https://www.ncbi.nlm.nih.gov/nuccore/NDIQ00000000.1/). Therefore, this emerging yeast is expected to greatly facilitate DCA production from fatty acid methyl esters, a vegetable oil-derived resource.

Wickerhamiella sorbophila was first isolated as a culture contaminant from washings of ion-exchange resins from a guanine monophosphate manufacturing plant [13]. Since its discovery, W. sorbophila has rarely been investigated, except for some short studies on its production of lactone [14] and chiral R-amino alcohol [15]. Here, we first described the development of effective genetic engineering tools for industrial application of W. sorbophila. To facilitate genetic engineering, selectable antibiotic-resistance markers were explored and used to remove target genes. To increase the probability of gene insertion into the target locus, the cell cycle of yeast was synchronized to the S/G2 phase, during which homologous recombination (HR) frequently occurs, for yeast transformation. Finally, we developed an engineered W. sorbophila strain with a blocked β-oxidation pathway and applied it to the biotransformation of DCA from methyl laurate.

Results and discussion

Cell growth in the presence of antibiotics

Genetic manipulation is essential for the development of industrially competent strains. Despite the fact that the newly isolated W. sorbophila strain was a strong candidate for DCA production, biological information for genetic engineering of this strain is lacking. As a basic step for the development of genetic engineering tools, antibiotics for use of selectable markers in W. sorbophila were investigated. We evaluated cell growth in the presence of several antibiotics that are primarily used in eukaryotic genetic engineering (Fig. 2). No inhibitory effect on cell growth was observed with blasticidin S and puromycin. Zeocin, hygromycin B, and G418 exerted a slight inhibitory effect, but cell growth was still observed even at high concentrations. However, nourseothricin
Fig. 1 The proposed biological pathway for dicarboxylic acid (DCA) production in Wickerhamiella sorbophila. Alkanes are first oxidized to the corresponding fatty alcohol by a cytochrome P450 monoxygenase/cytochrome P450 reductase complex (CYP + CPR). The fatty alcohol is converted into a corresponding fatty aldehyde by fatty alcohol oxidase (FAO) or alcohol dehydrogenase (ADH). Then, the fatty aldehyde is oxidized to fatty acid (FA) by fatty aldehyde dehydrogenase (FALDH). FA methyl ester is hydrolyzed by extracellular lipase or esterase before primary oxidation, and its corresponding FA is converted to DCA via the same mechanism as alkane. Converted FA and DCA are finally degraded to acetyl-CoA by the successive reactions of acyl-CoA synthetase (ACS), acyl-CoA oxidase (POX), 2-enoyl-CoA hydratase (MFE), 3-hydroxyacyl-CoA dehydrogenase (MFE), and 3-ketoacyl-CoA thiolase (POT). To prevent the continuous catabolism of carboxylic acids to acetyl-CoA, β-oxidation-related genes, POX, are removed.
Table 1 Studies investigating the biotransformation of medium-chain DCAs from alkanes, fatty acids, and fatty acid methyl esters

| Strain                  | Substrate       | Analytical method | DCA (g/l) | References |
|-------------------------|-----------------|-------------------|-----------|------------|
| **Petroleum-based resources** |                 |                   |           |            |
| C. tropicalis ATCC 20962 | Dodecane (C12)  | GC                | 140a      | [2]        |
| C. viswanathii ipe-1     | Dodecane (C12)  | pH titration      | 129.7     | [51]       |
| C. tropicalis CZ-15      | Tridecane (C13) | pH titration      | 98        | [3]        |
| C. tropicalis CGMCC 356  | Tridecane (C13) | pH titration      | 153       | [34]       |
| Y. lipolytica H222ΔPoF   | Dodecane (C12)  | GC                | 11        | [52]       |
| Y. lipolytica MTLY 37    | Dodecane (C12)  | GC                | 8         | [53]       |
| Y. lipolytica MTLY 37    | Tetradecane (C14) |  GC                | 2         | [53]       |
| **Vegetable oil-based resources** |                 |                   |           |            |
| C. tropicalis mutantb   | Methyl decanoate (C10) | GC             | 34.5      | [11]       |
| C. tropicalis ATCC 20962 | Methyl laurate (C12) | GC              | 66        | [35]       |
| C. tropicalis ATCC 20962 | Methyl myristate (C14) | GC             | 210b     | [2]        |
| C. cloacae FERM-P736    | Lauric acid (C12) | GC                | 5         | [54]       |
| W. sorbophila UHP4      | Methyl laurate (C12) | GC             | 92.5      | This work  |

* Concentration calculated based on the initial volume
b Strain tolerant to decanoic acid derived from C. tropicalis ATCC 20962

Fig. 2 Antibiotic sensitivity of W. sorbophila. Cell growth was examined at various concentrations of antibiotics to identify a selectable marker. Cells (1 × 10^7) were grown for 3 days on yeast peptone dextrose (YPD) agar medium containing blasticidin S, puromycin, Zeocin, hygromycin B, nourseothricin, or phleomycin. For comparison, cell growth is represented by “++” and “−−−” (+ ≤ 100 colonies; ++ ≤ 200 colonies; +++ ≤ 400 colonies; ++++ > 400 colonies).
and phleomycin totally inhibited cell growth at 100 and 200 μg/ml, respectively. *Streptomyces noursei nat1* and *Streptocalloteichus hindustanus ble* genes reportedly confer resistance to nourseothricin and phleomycin, respectively [16, 17]. These genes were consequently used as selectable markers for *W. sorbophila*.

**Heterologous expression of nat1 and ble genes**

Some yeasts, including the *Candida* species, decode CUG, a universal leucine codon, as serine. To enable heterologous expression of the marker genes, codon usage of *W. sorbophila* was analyzed. The analysis revealed that this strain might use standard genetic code (Additional file 1: Figure S1). Then, promoter and terminator sequences were investigated. While the use of heterologous promoter and terminator is often recommended to prevent interference caused by similarity of an endogenous DNA sequence during insertion of exogenous DNA into the chromosome, the stability or strength of expression is not guaranteed [18]. Hence, we next identified a homologous promoter sequence and a heterologous terminator sequence for heterologous gene expression. The 500-bp upstream region of the *TEF* gene was retrieved to identify a constitutive promoter in *W. sorbophila*. Translation elongation factor 1α gene, *TEF*, encoding the alpha subunit of the elongation factor complex, is involved in transferring the aminoacyl tRNA to the ribosome, and its promoter has been reported to constantly drive gene expression, regardless of the external environment and growth stage [19]. A heterologous terminator was obtained from the *C. tropicalis GAPDH* gene. To test the feasibility of using expression cassettes containing the *nat1* or *ble* gene, we transformed *W. sorbophila* with the constructs by allowing linearized DNA (the resistance cassettes) lacking homology arms to be randomly inserted into the chromosome (Fig. 3). After 4 days of incubation, nourseothricin- or phleomycin-resistant colonies appeared on the selection medium, and the inserted genes were identified by colony polymerase chain reaction (PCR) of the transformants. As shown in Fig. 3, the PCR products corresponding to the size of the inserted gene were detected in all resistant colonies tested, but not in the wild type. These observations indicated that the transcription system involving marker genes using a homologous promoter and heterologous terminator operated well in *W. sorbophila*.

**HR efficiency**

In eukaryotes, double-strand breaks (DSBs) induce two kinds of DNA repair mechanisms: HR and non-homologous end joining (NHEJ). HR allows the truncated 3’-end strand to recognize and replicate intact homologous double strands. NHEJ directly connects two DSBs by annealing short sequences near their end [20]. These two mechanisms occur competitively, and the balance varies depending on the cell cycle or species [21]. In yeast genetic engineering, HR has been used for insertion of external DNA at a desired position in the chromosome. Nonconventional yeasts, such as *Y. lipolytica* and *Kluyveromyces marxianus*, reportedly predominantly use NHEJ, resulting in random integration. To increase the HR rate in nonconventional yeast, most studies have focused on eliminating the function of the genes associated with the NHEJ pathway, such as the ones encoding Ku70/80 [22, 23] and DNA ligase IV [24]. However, the errors of the highly conserved DNA repair system impair the maintenance of genome integrity. Mutations in the DNA repair system increase the susceptibility to DNA damage, and the resulting increased mutation rate reduces cell viability [25, 26]. To maintain an intact DNA repair system, we increased the HR rate by regulating the cell cycle. Recent studies have shown that the balance of HR and NHEJ shifts during the cell cycle [20, 21] (Fig. 4a). In yeast and mammalian cells, since the accessibility of sister chromatids—the preferred template for HR repair—affects HR efficiency, HR use is elevated between the S and G2 phases of the cell cycle, during which the sister chromatids can be used [27]. Therefore, we deleted the target genes by synchronizing the cells in the S phase using HU, an inhibitor of ribonucleotide reductase [28], and compared the HR efficiency in the resultant cells with that attained by other existing methods.

Prior to this experiment, the appropriate concentration of HU was determined. Microscopic analysis (Fig. 4b) revealed that cells at the S/G2/M phase coexisted at 0.05 and 0.1 M of HU, whereas cell division was inhibited at 0.4 M. Therefore, HU was used at 0.2 M to arrest the cells at the S/G2 phase. To compare the HR efficiency depending on the cell growth conditions, deletion cassettes containing selectable markers were constructed and integrated into the *URA3, ADE2, POX1*, and *POX2* loci, accordingly, depending on the growth stage (Additional file 1: Figure S2) (Table 2). *ura3* or *ade2* was eliminated using the *ble* marker; the resultant strains were named *W. sorbophila* U1 (*ura3Δ*) and *W. sorbophila* A1 (*ade2Δ*). *pox1* or *pox2* was eliminated using the *URA3* marker; the resultant strains were named *W. sorbophila* UHP1 (*pox1Δ* with *URA3* harboring glu) and *W. sorbophila* UHP2 (*pox2Δ* with *URA3* harboring glu), respectively. In haploid yeast, the removal of genes with physiologically important functions is detrimental to cell growth and leads to a delay in the appearance of transformants [29]. To reduce the probability of selection errors, we supplemented the selection medium with the required nutrients, i.e., adenine and uracil, to support the growth of the desirable transformants. As shown in Table 2, HR
Fig. 3  Schematic representation of the expression cassettes and chromosome integration in W. sorbophila. The pUC18-nat1 (a) and pUC18-ble (b) cassettes, each containing the TEF promoter and GAPDH terminator, were PCR amplified using primers TEFp-F and GAPt-R and randomly inserted into the chromosome. The gel images show the results of yeast colony PCR using primers TEFp-F and GAPt-R. The sizes of the bands are 1.4 kb (a) and 1.2 kb (b).
frequency was highest in HU-treated cells, followed by those in the logarithmic and stationary phases of growth, in that order. HR efficiency at the POX1 and POX2 loci increased from 56.3% and 41.7%, respectively, to 97.9% for both cases. The original HR efficiency at the URA3 and ADE2 loci was nearly 0% in both early stationary and logarithmic phases, and increased to 4.8% and 25.6%, respectively, after HU treatment.

The experiment revealed some distinct features of W. sorbophila. The strain showed various HR efficiencies depending on the targeted gene locus. The HR efficiency of genes essential for growth, such as ADE2 and URA3, was close to 0%, but that of POX genes involved in fatty acid metabolism exceeded 40% (Table 2). This suggested the occurrence of HR hotspots on the genome of W. sorbophila. We successfully removed the ADE2 and URA3 genes, typically characterized by nearly 0% HR efficiency, by regulating the cell cycle. Therefore, transformation involving HU is an effective and easy approach of increasing HR efficiency without permanently damaging the DNA repair system.

Characterization of the putative WsPOX genes

Acyl-CoA oxidase (EC:1.3.3.6.), which catalyzes the initial step in fatty acid β-oxidation, transfers hydrogen atoms in the CH–CH group of acyl-CoA to molecular oxygen [30], resulting in the formation of trans-2-enoyl-CoA and H₂O₂. In alkane-assimilating yeasts, the deletion of POX genes prevents degradation of fatty acids, promoting ω-oxidation of the terminal methyl group [2]. There are six POX genes in Y. lipolytica, some of which exhibit specific substrate affinity, depending on the carbon chain length [31, 32]. Based on the whole-genome sequence of W. sorbophila, two putative POX genes with high amino acid sequence similarity to YALI0F10857g (the Y. lipolytica POX2 gene) were identified (POX1: B9G98_00908, 60% sequence similarity; POX2: B9G98_04160, 52% sequence similarity).

We characterized the putative POX genes by analyzing pox gene-deficient mutants. The pox1 and pox2 deletion strains were obtained using the methods described above. For the sequential disruption of the POX genes, the URA3 marker of W. sorbophila UHP1 was removed by HR between the both terminal glu sequences

Table 2  HR efficiency depending on the growth stage

| Growth stage | Strain | Target gene | Marker gene | HR efficiency |
|--------------|--------|-------------|-------------|---------------|
| Early stationary phase | DS02 | ADE2 | ble | 0% (1365)a |
| | DS02 | URA3 | ble | 0% (186)a |
| | UHP1 | POX1 | URA3 | 56.3% (48)a |
| | UHP2 | POX2 | URA3 | 41.7% (48)a |
| Log phase | DS02 | ADE2 | ble | 1% (1470)a |
| | DS02 | URA3 | ble | 0% (186)a |
| | UHP1 | POX1 | URA3 | 81.3% (48)a |
| | UHP2 | POX2 | URA3 | 83.3% (48)a |
| S arrest | DS02 | ADE2 | ble | 25.6% (379)a |
| | DS02 | URA3 | ble | 4.8% (186)a |
| | UHP1 | POX1 | URA3 | 97.9% (48)a |
| | UHP2 | POX2 | URA3 | 97.9% (48)a |

* Number of transformants screened
(pop-out) \((W.\ sorbophila\ UHP3,\ \textit{pox}1\Delta\ \textit{without\ \textbf{URA}3\ harboring\ glu})\), and then the \textit{POX2} gene was eliminated by reusing the \textit{URA3} marker \((W.\ sorbophila\ UHP4,\ \textit{pox}1\Delta/\textit{pox}2\Delta\ \textit{with\ \textbf{URA}3\ harboring\ glu})\). As shown in Fig. 5a, b, the overall growth of mutants was reduced in comparison with that of the wild-type strain. The \textit{W. sorbophila UHP2} strain grew in the presence of all tested carbon sources. However, the \textit{W. sorbophila UHP1} strain exhibited unaltered growth on glucose, but not in the presence of medium-chain \((C_{10−C_{12}})\) and long-chain \((C_{14−C_{18}})\) alkanes. Only the \textit{POX1} gene—with a broad substrate spectrum—is expected to play an important role in fatty acid metabolism. An interesting phenomenon was reported previously, namely, that \textit{W. sorbophila} simultaneously consumes various substrates during culture in the presence of mixed fatty acid methyl esters \((C_{10−C_{16}})\) \cite{12}. This may be associated with the broad substrate specificity of the product of the \textit{POX1} gene.

The \textit{POX2} gene, however, is considered to be a pseudo-gene or involved in the metabolism of short-chain fatty acids. The DDDA conversion test (Fig. 5c) revealed that only \textit{W. sorbophila UHP1} and \textit{W. sorbophila UHP4} were able to convert dodecane to DDDA. \textit{W. sorbophila UHP1} and \textit{W. sorbophila UHP4} produced 9.38 and 9.12 g/l of DDDA, respectively, and their conversion yields \((\text{mol/mol})\) were 90.1% and 92.6%, respectively. By contrast, the growth of \textit{W. sorbophila UHP2} was reduced during the first 24 h, similarly to the other mutants, but eventually metabolized and used dodecane for cell growth. Hence, removal of the \textit{POX1} gene is essential for the accumulation of DDDA.

### Fed-batch fermentation for DDDA production

In \textit{C. tropicalis}, a well-known DCA-producing yeast, the fermentation process for DCA production has already been optimized \cite{2, 33, 34}. The fermentation for DCA production is divided into growth and conversion stages, where the medium pH greatly affects fatty acid and DCA solubility, DCA excretion and cell viability \cite{11, 34}. During the conversion stage, acidic conditions (=pH 5.8) steadily promoted the cell growth, but DCA precipitates were formed and interfered with the process \cite{35}. In the resting cells, a greater amount of DCA was released at a higher pH \cite{36}; however, in living cells, pH values above 8.2 reduced cell viability and DCA productivity \cite{34}. Considering this, we performed fed-batch fermentation using \textit{W. sorbophila} UHP4 (with completely blocked \(\beta\)-oxidation) and methyl laurate. The pH was adjusted to 5.5 for the initial 12 h period to obtain high cell density, after which the pH was increased to 7.2–8.0.
to solubilize fatty acid and DCA, and to promote DCA secretion (as described in detail in “Methods”).

DCA conversion is initiated by cytochrome P450 monooxygenase (CYP); members of the CYP family are alkane-inducible in alkane-assimilating yeasts [37–39]. In C. tropicalis, the addition of methyl decanoate without the pre-activation of ω-oxidation resulted in the accumulation of decanoic acid and directly affected cell viability [11]. Thus, prior to methyl laurate addition, 1% (v/v) dodecane of the culture was added to pre-activate the ω-oxidation pathway of W. sorbophila. In addition, Tween 80, a nonionic surfactant esterified with polyethoxylated sorbitan and oleic acid, was used to facilitate the uptake of dodecane. In various prokaryotic and eukaryotic microorganisms, cellular uptake of hydrophobic substrates is involved in several mechanisms, such as substrate solubilization by extracellular emulsifying agents, modification of the cell surface to facilitate adhesion [40], passive diffusion via the cell membrane [33], and transporter-mediated uptake [41]. In C. tropicalis, the alkane-binding affinity of glucose-grown cells was lower than that of alkane-grown cells due to the alteration of the cell wall components [42]; the uptake of alkane on glucose-grown cells showed a greater dependence on the emulsification of alkane than alkane-grown cells [42]. Based on these reports, 1% (v/v) Tween 80 was added to dissolve dodecane in the culture medium. Even if Tween 80 was hydrolyzed, as observed in some yeasts [43], oleic acid resulting from Tween 80 cleavage is not expected to affect the amount of DDDA produced by the strain with blocked β-oxidation. Finally, W. sorbophila UHP4 produced 92.5 g/l of DDDA over 126 h, with the productivity and yield of 0.83 g/l/h and 0.86 (mol/mol), respectively (Fig. 6b). This was 1.4 times higher than the fermentation data for C. tropicalis and methyl laurate (Table 1).

Most studies with dodecane have reported higher values than those obtained in the current study; however, an accurate comparison was difficult because the DDDA concentration was determined by pH titration or calculated based on the initial volume (Table 1). In the absence of Tween 80, DDDA productivity was greatly reduced, almost by half (Fig. 6a). A detailed study of this phenomenon would aid the improvement of the W. sorbophila performance. Furthermore, alongside process optimization, future studies on biotransformation using fatty acid methyl esters with various carbon chain lengths would expand the scope for the application of this strain.

Conclusions

Currently, sustainable production of DCA is a major concern in industrial biotechnology. Although many studies have contributed to this field, the biotransformation of DCA from renewable resources has been hindered because of the cellular toxicity of fatty acids. Previously, we identified several positive aspects of DCA production—industrially and biologically—using a W. sorbophila strain [12]. Here, we developed a novel microbial platform for DCA production, overcoming the limitation of toxicity. Based on whole-genome sequence information for W. sorbophila, we developed and optimized genetic engineering tools. To increase HR efficiency during the yeast transformation process, the cell cycle was artificially fixed at the S/G2 phase. To prevent the metabolism of alkanes and fatty acids, putative POX genes were characterized and removed. Interestingly, only pox1-deficient strains produced DDDA, suggesting that the POX1 gene plays a major role in β-oxidation. In 5-l fed-batch fermentation with W. sorbophila UHP4, 92.5 g/l of DDDA was produced from methyl laurate in 126 h. In conclusion, the presented data demonstrated that W. sorbophila is a promising microbial platform for DCA production from vegetable oil-based resources. Furthermore, the developed genetic engineering tools will form the basis for future research for the production of various value-added oleochemicals derived from vegetable oil, together with traditional metabolic engineering strategies, to improve DCA productivity.

Methods

Strains, materials, and growth conditions

The strains used in the current study are listed in Table 3. The nat1 and ble genes were synthesized by GenScript (Piscataway, NJ, USA) (Additional file 1: Table S1). YNB medium without amino acids, and yeast extract and peptide were purchased from Becton-Dickinson (Franklin Lakes, NJ, USA). Nourseothricin was purchased from BioVision (Milpitas, CA, USA). Blasticidin S, phleomycin, puromycin, and Zeocin were purchased from InvivoGen (San Diego, CA, USA). Adenine, uracil, HU, alkanes, lauric acid, methyl laurate, DDDA, Tween 80, and N,O-bis-trimethylsilyl-trifluoroacetamide were from Sigma-Aldrich (St. Louis, MO, USA). 5-Fluoroorotic acid (5-FOA) and tetradecanedioic acid were from TCI Chemical (Tokyo, Japan). Antifoam agent (adekanol LG-109) was from Adeka (Tokyo, Japan). All other chemicals were purchased from Junsei Chemical (Tokyo, Japan).

The yeast strains were routinely cultured at 30 °C in YPD medium. When required, the medium was supplemented with 100 mg/l adenine or 100 mg/l uracil. For
the selection of antibiotic markers, $1 \times 10^7$ cells were spread on YPD agar medium containing 50–300 μg/ml blasticidin S, 10–40 μg/ml puromycin, 100–400 μg/ml Zeocin, 100–600 μg/ml hygromycin B, 1–4 mg/ml G418, 50–300 μg/ml nourseothricin, or 100–400 μg/ml phleomycin. Antibiotic-resistant colonies were observed after incubation at 30 °C for 3 days.

To analyze the growth of W. sorbophila strains, appropriate carbon sources [20 g/l glucose, 1% (v/v) decane, 1% (v/v) dodecane, 1% (v/v) tetradecane, 1% (v/v) hexadecane, or 1% (v/v) octadecane] were added to YNB medium with 0.1% (v/v) Tween 80. In the case of solid YNB medium, alkanes were supplied in the vapor phase, as described previously [44]. Alkanes with fewer than 8
or over 20 carbons were excluded because of their toxicity, volatility, or melting temperature. Pre-cultured cells were washed with YNB medium and then spotted onto agar medium in tenfold serial dilutions starting with 10 μl of suspensions with OD₆₀₀ of 0.1.

**Prediction of codon usage**
The CUG codon usage of *W. sorbophila* was predicted using the fungal CUG codon-predicting site Bagheera [45]. The whole-genome sequence information was obtained from the GenBank database (Accession Number CP020102.1).

**PCR amplification**
The *nat1* and *ble* genes were PCR amplified using primers *nat1*-F and *nat1*-R, and *ble*-F and *ble*-R, respectively. A 500-bp sequence upstream of the *TEF* gene was amplified for use as a promoter from the genomic DNA of *W. sorbophila* DS02 using primers TEFp-F and TEFp-R. The 500-bp 5′ and 3′ homologous arms of the *ADE2* gene were amplified for target gene disruption from the genomic DNA of *W. sorbophila* DS02 using primers ADE2 HR-F1, ADE2 HR-R2, ADE2 HR-F3, and ADE2 HR0-R4, for *ADE2* disruption; *URA3* HR-F1, *URA3* HR-R2, *URA3* HR-F3, and *URA3* HR-R4, for *URA3* disruption; *POX1* infu-F1, *POX1* infu-R2, *POX1* infu-F3, and *POX1* infu-R4, for *POX1* disruption; and *POX2* infu-F1, *POX2* infu-R2, *POX2* infu-F3, and *POX2* infu-R4, for *POX2* disruption. The genomic sequence information was obtained from the GenBank database (Accession Number CP020102.1). A 300-bp fragment downstream of the *GAPDH* gene sequence was amplified for use as a terminator from the genomic DNA of strain ATCC 20962 using primers CtGAPt-F and CtGAPt-R; the genomic sequence information was obtained from the GenBank database (Accession Number CP020102.1).

**Table 3 Strains used in the current study**

| Strain            | Genotype                        | Reference |
|-------------------|---------------------------------|-----------|
| *W. sorbophila* DS02 | Wild type                       | [12]      |
| *W. sorbophila* U1  | *ura3Δ::Sable*                   | This work |
| *W. sorbophila* A1  | *ade2Δ::Sable*                   | This work |
| *W. sorbophila* UHP1 | *ura3Δ::pox1Δ::glu-CtURA3::glu* | This work |
| *W. sorbophila* UHP2 | *ura3Δ::pox2Δ::glu-CtURA3::glu* | This work |
| *W. sorbophila* UHP3 | *ura3Δ::pox1Δ::glu*             | This work |
| *W. sorbophila* UHP4 | *ura3Δ::pox1Δ::glu pox2Δ::CtURA3* | This work |

**Plasmid construction**
To construct a heterologous gene expression cassette, the *TEF* promoter, *nat1* gene, and *GAPDH* gene were amplified for use as a promoter from the genomic DNA of *W. sorbophila* DS02 using primers TEFp-F and TEFp-R. Disruption of target genes was first confirmed by colony PCR, and then by PCR with the genomic DNA isolated from the transformants, using primers *URA3con*-F, *URA3con*-R, *POX1con*-F, *POX2con*-F, and *POXcon*-R, as appropriate, specific for the sequence outside of the homologous arms. All primers used in the current study are listed in Additional file 1: Table S2.

**Transformation**
*Wickerhamiella sorbophila* was transformed using the lithium acetate method, with some modification [47]. After transformation of cells with 1 μg of the linearized cassette, the cell pellet was suspended in 1 ml of YPD and incubated for 2 h at 30 °C with shaking at 180 rpm. When
required, the medium was supplemented with 100 mg/l adenine or 100 mg/l uracil. Then, the cells were spread on the selection medium and incubated for 5 days at 30 °C. Ura3-deletion candidates were confirmed by colony PCR after primary counter-selection on 5-FOA agar medium [48]; ade2-deletion candidates were simply identified by red/white colony color [49]. Putative pox gene-deletion candidates were confirmed by colony PCR.

Cell synchronization using HU
Synchroonization of W. sorbophila cells was performed according to a previously published protocol [50]. Subsequently, subcultured cells were grown to OD600 = 1 in 50 ml of YPD medium (when required, 100 mg/l adenine or 100 mg/l uracil was included in the medium), treated 50 ml of YPD medium (when required, 100 mg/l adenine or 100 mg/l uracil was included in the medium), treated with HU, and incubated for 2 h at 30 °C with shaking at 180 rpm. The synchronized cells were used for yeast transformation.

Marker recycling
To remove the URA3 marker from the W. sorbophila UHP1 genome, the cells were grown in 3 ml of YPD medium overnight. Further, 100 μl of the culture was spread on 5-FOA agar medium and incubated for 5 days at 30 °C. The identity of the obtained colonies was confirmed by colony PCR.

Flask cultivation for DCA production
One loopful of cells was pre-cultured in a 15-ml round tube with 3 ml of YPD medium overnight. Then, 1 ml of the culture was transferred to a 250-ml baffled flask containing 20 ml of YPDP medium (10 g/l yeast extract, 20 g/l peptone, 30 g/l glucose, and 1.8 g/l KH2PO4) and incubated at 30 °C with shaking at 200 rpm. After culturing for 24 h, 0.3 g (15 g/l) K2HPO4 and 0.2 ml (1%, v/v) of dodecane were added to the culture broth; 0.1 g (5 g/l) glucose was continuously added at a rate of 2 g/l/h; the medium pH was adjusted from 5.5 to 7.2 with 8 N NaOH, and then increased up to 7.8 (pH 7.2 for the culture period of 12–24 h; pH 7.4 for 24–48 h; pH 7.6 for 48–96 h; pH 7.8 for 96–126 h; and pH 8.0 for 120–126 h). After 24 h, methyl laurate was continuously added at a rate of 0.5–2 ml/l/h (0.5 ml/l/h for the culture period of 24–26 h; 1 ml/l/h for 26–30 h; 1.2 ml/l/h for 30–42 h; 1.6 ml/l/h for 42–72 h; 1.8 ml/l/h for 72–96 h; and 2 ml/l/h for 96–126 h) (Fig. 6b). The feeding rate was calculated based on the initial volume, and product concentration and productivity were calculated based on the end volume.

Fed-batch fermentation
Fed-batch fermentation using the engineered W. sorbophila strain was performed according to a previously described protocol, with some modification [12]. One loopful of cells grown on YPD agar plates was inoculated in a 250-ml baffled flask containing 20 ml of YPDP medium and incubated at 30 °C with shaking at 200 rpm for 24 h. The culture was transferred to a 2-l baffled flask containing 200 ml of YPD medium and incubated at 30 °C and 200 rpm for 24 h. Subcultured cells were transferred to a 5-l fermenter (CNS, Daejeon, Korea) with 2 l of medium containing 50 g/l glucose, 20 g/l yeast extract, 8 g/l (NH4)2SO4, 2 g/l K2HPO4, 0.1 g/l NaCl, 0.1 g/l CaCl2 · 2H2O, and 0.5 ml of antifoam agent. Temperature, medium pH, and aeration were controlled at 30 °C, 5.5, and 1 vvm, respectively. Dissolved oxygen was maintained above 30% by modulating the agitation speed between 200 and 900 rpm. After culturing for 12 h, 20 ml (1%, v/v) of dodecane and 2 ml (0.1%, v/v) of Tween 80 were added to the culture broth; glucose was continuously added at a rate of 2 g/l/h; the medium pH was adjusted from 5.5 to 7.2 with 8 N NaOH, and then increased up to 7.8 (pH 7.2 for the culture period of 12–24 h; pH 7.4 for 24–48 h; pH 7.6 for 48–96 h; pH 7.8 for 96–126 h; and pH 8.0 for 120–126 h). After 24 h, methyl laurate was continuously added at a rate of 0.5–2 ml/l/h (0.5 ml/l/h for the culture period of 24–26 h; 1 ml/l/h for 26–30 h; 1.2 ml/l/h for 30–42 h; 1.6 ml/l/h for 42–72 h; 1.8 ml/l/h for 72–96 h; and 2 ml/l/h for 96–126 h) (Fig. 6b). The feeding rate was calculated based on the initial volume, and product concentration and productivity were calculated based on the end volume.

Analytical methods
Cell density, the concentrations of glucose, and DDDA in the culture broth were determined. Cell density was determined using an ultraviolet spectrophotometer (Uvikon XL; Secomam, Alès, France) by measuring the absorbance at 600 nm. Glucose concentration was determined using a glucose analyzer (YSI 2700; YSI Life Sciences, Yellow Springs, OH, USA). DDDA was analyzed by gas chromatograph (Master GC; Dani Instruments, Cologno Monzese, Italy) equipped with an RTX-5 column (Restek, Bellefonte, PA, USA). A 50-μl aliquot of 6 N H2SO4 was added to 100 μl of culture broth, with 0.1 N NaOH, and then increased up to 7.8 (pH 7.2 for the culture period of 12–24 h; pH 7.4 for 24–48 h; pH 7.6 for 48–96 h; pH 7.8 for 96–126 h; and pH 8.0 for 120–126 h). After 24 h, methyl laurate was continuously added at a rate of 0.5–2 ml/l/h (0.5 ml/l/h for the culture period of 24–26 h; 1 ml/l/h for 26–30 h; 1.2 ml/l/h for 30–42 h; 1.6 ml/l/h for 42–72 h; 1.8 ml/l/h for 72–96 h; and 2 ml/l/h for 96–126 h) (Fig. 6b). The feeding rate was calculated based on the initial volume, and product concentration and productivity were calculated based on the end volume.

Additional file 1: Table S1. The sequences of nat1 and ble genes used in the current study. Table S2. Primers used in the current study. Figure S1. Schematic representation of the deletion cassettes and chromosome integration in W. sorbophila. The genome sequence uploaded in Bagheera was matched with sequences of 2071 proteins from 38 different protein families using TBLASTN [1]. Figure S2. Schematic representation of the deletion cassettes and chromosome integration in W. sorbophila. The pUC18-URA3-ble cassette was PCR-amplified using primers URA3-F and URA3-R, and inserted into the URA3 locus (a). The pUC18-ADE2-ble cassette was PCR-amplified using primers ADE2-HF and ADE2-HR, and inserted into the ADE2 locus (b). The pGEM-POX1-que cassette was linearized by EcoRI digestion and inserted into the POX1 loci (c). The pGEM-POX2-que cassette was linearized by Smal digestion and inserted into the POX2 loci (d).
Abbreviations
S-FOA: 5-fluoroorotic acid; CYP: cytochrome P450 monoxygenase; DCA: dicarboxylic acid; DDDA: dodecanedioic acid; DS8: double-strand break; GC: gas chromatography; HR: homologous recombination; HU: hydroxyurea; NHE: non-homologous end joining; PA: polyamide; PCR: polymerase chain reaction; YNB: yeast nitrogen base; YPD: yeast peptone dextrose.

Authors' contributions
HJ and JA designed the entire research project; HL, CH, HWL, and GP executed the experimental work; HL wrote up the manuscript; HJ and WJ reviewed the manuscript; HL, revised the manuscript. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

Availability of data and materials
All datasets supporting the conclusions of this study are included in this published article (and its Additional file 1).

Consent for publication
Not applicable.

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Not applicable.

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