Role of Operon aaoSo-mutT in Antioxidant Defense in *Streptococcus oligofermentans*

Peng Zhou, Lei Liu, Huichun Tong*, Xiuzhu Dong*
State Key Laboratory of Microbial Resources, Institute of Microbiology, Chinese Academy of Sciences, Beijing, China

Abstract

Previously, we have found that an insertional inactivation of *aaoSo*, a gene encoding L-amino acid oxidase (LAAO), causes marked repression of the growth of *Streptococcus oligofermentans*. Here, we found that *aaoSo* and *mutT*, a homolog of pyrophosphohydrolase gene of *Escherichia coli*, constituted an operon. Deletion of either gene did not impair the growth of *S. oligofermentans*, but double deletion of both *aaoSo* and *mutT* was lethal. Quantitative PCR showed that the transcript abundance of *mutT* was reduced for 13-fold in the *aaoSo* insertional mutant, indicating that gene polarity derived from the inactivation of *aaoSo* attenuated the expression of *mutT*. Enzymatic assays were conducted to determine the biochemical functions of LAAO and MutT of *S. oligofermentans*. The results indicated that LAAO functioned as an aminoacetone oxidase [47.75 nmol H$_2$O$_2$ (min-mg protein)$^{-1}$]; and MutT showed the pyrophosphohydrolase activity, which removed mutagens such as 8-oxo-dGTP. Like paraquat, *aaoSo* mutations increased the expression of SOD, and addition of aminoacetone (final concentration, 5 mM) decreased the mutant’s growth by 11%, indicating that the *aaoSo* mutants are under ROS stress. HPLC did reveal elevated levels of cytoplasmic aminoacetone in both the deletion and insertional gene mutants of *aaoSo*. Electron spin resonance spectroscopy showed increased hydroxyl radicals in both types of *aaoSo* mutant. This demonstrated that inactivation of *aaoSo* caused the elevation of the prooxidant aminoacetone, resulting in the cellular ROS stress. Our study indicates that the presence of both LAAO and MutT can prevent endogenous metabolites-generated ROS and mutagens. In this way, we were able to determine the role of the *aaoSo-mutT* operon in antioxidant defense in *S. oligofermentans*.

Introduction

Oxidative stress is encountered universally by all organisms living in an oxygen environment. Oxygen is a relatively inert molecule in nature. However, when it is channeled to electron-transfer reactions, especially inside the cells that experience auto-oxidation of reduced enzyme cofactors, oxygen is converted into deleterious reactive oxygen species (ROS). Due to the higher energy that results from the presence of unpaired valence-shell electrons, ROS can cause severe damage to various biological macromolecules, especially DNA. This leads to mutagenesis, tumorogenesis, and aging [1,2].

Organisms have developed various means of resisting oxidative stress. Aerobes generally employ superoxide dismutase (SOD) and catalase cascades to eliminate cellular ROS. SOD dismutates superoxide anions as H$_2$O$_2$, which is then converted to H$_2$O and O$_2$ by catalase [3,4]. Anaerobes use the superoxide reductase (SOR)-dependent oxide decomposition pathway [5]. SOR reduces superoxide to H$_2$O$_2$ using reduced rubredoxin (Rd$_{red}$) as the electron donor. Then, in concert with the non-heme peroxidase ruberythrin, SOR reduces H$_2$O$_2$ to water. Intact H$_2$O$_2$ is not particularly deleterious to cells, but H$_2$O$_2$-derived hydroxyl radicals are indiscriminately damaging to the biological macromolecules. The conversion of H$_2$O$_2$ into radicals takes place through Fenton chemistry, which is activated by ferrous or copper ions [6–8]. Streptococci, which are facultative anaerobes, undergo fermentative metabolism to produce energy reserves. Streptococci do not possess catalase; instead, they have SOD and other oxidases. The formation of H$_2$O$_2$ is their main means of attenuating ROS damage [4,6]. *Streptococcus oligofermentans* is isolated from non-caries dental plaques in humans. It uses multiple pathways for the generation of H$_2$O$_2$ from oxygen [9]. These pathways include lactate oxidase (Lox) in oxidation of lactate to produce H$_2$O$_2$ [10]; L-amino acid oxidase (LAAO) to oxidize amino acids to H$_2$O$_2$ [11]; and a recently identified pyruvate oxidase (Pox) [12]. Lox and Pox contribute the majority of the H$_2$O$_2$ derived from glycolysis products [12], and LAAO generates lower levels of H$_2$O$_2$ when proteinaceous substances are present. Among streptococci, *S. oligofermentans* is one of those with the highest levels of tolerance to H$_2$O$_2$ [13]. This makes it a suitable model organism for the study of the mechanisms by which bacteria protect themselves against oxidative stress.

In a previous study on LAAO, we found that insertional inactivation of *aaoSo* caused severe reductions in the growth of *S. oligofermentans* [11]. To gain insight into the growth retardation, we performed a microarray assay and observed the differential expression of some genes in *aaoSo* mutants, including the genes encoding manganese-dependent SOD (*sodA*), DNA-binding ferritin-like protein, DNA repair enzyme, and NADH oxidase II (*nox*). Elevated H$_2$O$_2$ yield was detected in glucose-containing cultures of *aaoSo* mutants. These results imply that inactivation of *aaoSo* causes an elevation of cellular ROS levels in *S. oligofermentans*.
In the present study, we used genetic and biochemical analyses to find that LAAO is an authentic aminoacetone oxidase. It reduces the aminoacetone-derived ROS in vivo. Furthermore, aaoSo, by constituting an operon with mutT, encoding a nucleotide-oxide scavenging protein, develops a dual guard by reducing the production of cellular ROS and clearing the nucleotide oxides generated by ROS in streptococci. Thus, a novel antioxidant defense gene operon has been demonstrated.

**Results**

**Identification of the aaoSo-mutT Operon in S. oligofermentans**

Previous studies have found that an insertional mutation of aaoSo decreased the growth of S. oligofermentans to two thirds that of the wild-type strain [11]. This suggests that other inhibitory mechanisms might be present in addition to the H2O2 generated by LAAO. To evaluate possible inhibitive causes, we examined the genes that flanked aaoSo in the genome of S. oligofermentans. As shown in Fig. 1A, a putative gene encoding MutT (Acc No. JQ083601), a homolog of pyrophosphohydrolase (8-oxo-dGTPase) in *Escherichia coli* [14], was located 19 bp downstream of aaoSo. The two appear to constitute an operon. Then, using a reverse transcription-PCR assay, we determined that aaoSo and mutT were cotranscribed (Fig. 1B). This confirmed the presence of the operon and coexpression of the two genes in S. oligofermentans.

**Role of Coexistence of aaoSo and mutT in S. oligofermentans**

To determine the relative effects of aaoSo and mutT on the growth of S. oligofermentans, we constructed deletion mutants for each gene. As shown in Fig. S1, the deletion mutant ΔaaoSo exhibited slightly slower growth than wild-type strain, and ΔmutT did not show any detectable decrease in growth. A complementary strain of the insertional mutant of aaoSo resumed growth (Fig. 2). However, we never obtained a double gene mutant, here denoted ΔaaoSo::ΔmutT, even under obligate anaerobic conditions by manipulating inside an anaerobic chamber. This implies that double mutation of the two genes may be lethal. To confirm this, we first cloned the aaoSo-mutT operon into the *E. coli*-streptococcus shuttle-expression vector pDL278 and then introduced it into the wild-type strain of S. oligofermentans. Under this background, we readily deleted aaoSo::mutT from the chromosome. This mutant also carried a plasmid containing the operon, which showed growth patterns similar to the wild-type strain. To clarify the effect of an insertional mutation of aaoSo on the downstream mutT, we determined the expression of mutT by quantitative PCR and found a 13-fold reduction in transcript abundance relative to that of the wild-type strain, but in the ΔaaoSo strain, the deletion mutant, mutT expression was similar to the wild-type (Table 1). This indicates that aaoSo inactivation caused a gene-polarity effect on mutT. This insertional mutant was designated ΔaaoSo-mutT, and the suppression of growth was predicted to be related to the loss of MutT activity. Taken together, the coexistence of aaoSo and mutT

![Figure 1. Cotranscription of aaoSo and mutT in S. oligofermentans.](image-url)
Methylglyoxal (MG) and H2O2 [15,16]. Given that aminoacetone has been shown to remove aminoacetone by converting it to semicarbazide-sensitive amine oxidase (SSAO) has been identified as an electron donor in reactions involving active transition metals. Aminoacetone belongs to the amine family, and is determined as (data not shown).

Figure 2. Growth of the wild-type, aaoSo-mutT mutant of S. oligofermentans and its complementary strain in glucose. Strains were pre-cultured overnight in TYG medium, and 1:100 diluted with fresh TYG medium with the same OD600. Samples were taken every 1 h to detect OD600. Symbols: △, wild-type strain; ○, aaoSo-mutT mutant; ◆, aaoSo-mutT complementary strain. The results are shown as the means±SD of three independent experiments.

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seems to be essential for S. oligofermentans to carry out its normal physiological functions and even survive.

LAAO Authentically Acts as an Aminoacetone Oxidase

Enzymatic activity was assayed to clarify the function of LAAO. Our previous study showed that LAAO, an L-amino acid oxidase, mainly oxidizes seven kinds of amino acids to generate H2O2 [11]. Aminoacetone belongs to the amine family, and is determined as an electron donor in reactions involving active transition metals. So that it generates superoxide anions (ROS) from oxygen [15]. In human cells, semicarbazide-sensitive amine oxidase (SSAO) has been shown to remove aminoacetone by converting it to methylglyoxal (MG) and H2O2 [15,16]. Given that aminoacetone is an analog of amino acids, we tested the enzymatic activity of the overexpressed LAAO on aminoacetone. By using 25 mM of aminoacetone as substrate and air as an oxygen supply, we detected the production of 47.75 nmol H2O2 (min·mg protein)−1. In parallel, the activity of LAAO on lysine was only 2.22 nmol H2O2 (min·mg protein)−1. Even when lysine concentration was increased to 100 mM, the activity was only 12.29 nmol H2O2 (min·mg protein)−1. Similar low activities were detected for aspartic acid and glutamine (Fig. S2). This indicates that aminoacetone, not amino acids, is the preferred substrate of LAAO. However, unlike the described amine oxidases [17], LAAO did not catalyze benzylamine or methylamine conversion (data not shown).

Table 1. Transcript levels of mutT and mutX in the wild-type, aaoSo, and aaoSo-mutT mutants of S. oligofermentans.

| Strains      | The number of gene transcript* |
|--------------|-------------------------------|
|              | mutT                          | mutX                         |
| wild-type    | 0.66±0.15                     | 0.03±0.01                    |
| aaoSo-mutT   | 0.05±0.01                     | ND                           |
| aaoSo        | 0.76±0.17                     | ND                           |

*The transcript copies are given as the means±SD/10,000 16S rRNA copies. ND, not determined.

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Role of MutT in Elimination of Nucleotide Oxides in S. oligofermentans

Previous research has shown that dGTP can be readily oxidized to 8-oxo-dGTP by hydroxyl radicals, owing to its hypersensitivity to oxidation [18,19]. MutT of E. coli and its human homolog hMTH-1 protein hydrolyze 8-oxo-dGTP to pyrophosphate (PPi) and the harmless monophosphate 8-oxo-dGMP through a specific pyrophosphohydrolase (8-oxo-dGTPase) activity. Two mutT homologs, mutT and mutX, were found in the genome of S. oligofermentans. By using quantitative PCR, we determined that the transcript abundance of mutT was about 20-fold higher than that of mutX (Table 1), similar to the results of our previous microarray assays (unpublished data).

Next, we determined the pyrophosphohydrolase activity of the purified MutT protein of S. oligofermentans in vitro. According to the activity of MutT of E. coli, synthetic 8-oxo-dGTP was used as the substrate in the enzymatic assay. Upon addition of the purified recombinant MutT protein from S. oligofermentans, reduction of 8-oxo-dGTP levels was detected in the reaction mixture by HPLC (Fig. S3). This confirmed the pyrophosphohydrolase activity of S. oligofermentans MutT.

Effects of Inactivation of aaoSo on the Regulation of sodA

Our previous microarray data identified increased mRNA from sodA, the only SOD-encoding gene in the S. oligofermentans genome, by the aaoSo mutant relative to the wild-type strain (unpublished data). As superoxide anion is the well-defined substrate of SOD, a link between LAAO protein and the cellular ROS level may exist. To test this hypothesis, quantitative PCR was used to assay the expression of sodA in the aaoSo mutants of S. oligofermentans. As expected, higher transcriptional abundance of sodA was detected in aaoSo-mutT mutant (76.63±13.62 copies) and aaoSo mutant (47.94±8.03 copies), relative to the wild-type strain (13.11±3.02 copies). In addition, dramatic increases in Mn-SOD activity were observed in the early and middle log phase cultures of both mutants (Table 2). As sodA mRNA levels increase 3-fold in wild-type S. oligofermentans treated with the redox-cycling drug paraquat, which induces production of cellular endogenous superoxides (data not shown), we propose that the accumulation of superoxide anion increases the expression of Mn-SOD in aaoSo-mutT and aaoSo mutants.

We then constructed AaoSo and AaoSo-mutT-AaoSo mutants to confirm the superoxide-induced impairment of growth. As expected, no obvious decrease in growth was observed for AaoSo mutant. However, a reduction in growth of nearly 70% was detected for the double mutant, AaoSo-mutT-AaoSo (Fig. S4). Whereas when grew under strictly anaerobic conditions, AaoSo-mutT mutant grew to a final OD600 of 1.2, similar to wild-type; and the growth of the mutant AaoSo-mutT-AaoSo was restored to a level similar to that of the insertional mutant of aaoSo. This implies that superoxide anions may be the key to impairing the growth of AaoSo-mutT mutants.

Table 2. Mn-SOD activities of deletion and insertion mutants of S. oligofermentans aaoSo.

| Growth time | Specific activity (U mg−1 protein)* |
|-------------|------------------------------------|
|             | aaoSo                             | aaoSo-mutT                         |
| 4 h         | 2.55±0.68                         | 34.89±4.92                         |
| 6 h         | 37.6±5.33                         | 45.62±8.68                         |

*Activity was detected using the method of pyrogallol auto-oxidation [40]. Data were obtained as means±SD from three independent experiments.

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Figure 3. ESR spectra of the DMPO treated cells of *S. oligofermentans*. The characteristic 1:2:2:1 quartet of hydroxyl radical adduct of DMPO signals were shown in the DMPO-contained cell suspensions of (A) *aaoSo-mutT* mutant; (B) *aaoSo* mutant, and (C) wild-type strain.

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**Effects of Mutation of *aaoSo* on Cellular Levels of ROS**

To further evaluate the link between intracellular levels of ROS to LAAO, we measured the ROS levels in the *aaoSo-mutT*, *aaoSo* mutants and the wild-type strain. Considering that the accumulation of superoxide in cells is concomitant with abundant H2O2 generated by lactate oxidase and pyruvate oxidase, and either superoxide anion or H2O2 can cause the release of iron from proteins containing [4Fe–4S]2+ clusters, the Fenton reaction can cause an elevation in cellular hydroxyl radicals [20]. Therefore, hydroxyl radicals were assessed in *S. oligofermentans* strains using electron spin resonance (ESR) with 5,5-dimethyl-1-pyrroline-N-oxide (DMPO) spin trap [21,22]. ESR spectroscopy is the most efficient and non-destructive technique for detecting chemical species that have unpaired electrons, such as the superoxide anion radical and hydroxyl radical. As a spin trap, DMPO has been frequently used to detect oxygen-centered radicals [22]. As shown in Fig. 3, intense and characteristic 1:2:2:1 quartet of DMPO-OH• adduct signal was obtained both in the *aaoSo-mutT* and *aaoSo* mutants, but only a very weak signal was detected in the wild-type strain. This demonstrates that the *aaoSo* mutants were under a relatively high ROS stress.

**Effects of Mutation of *aaoSo* on Levels of Cytoplasmic Aminoacetone**

To evaluate the link between intracellular ROS and the inactivation of LAAO, based on the aminoacetone oxidase activity of LAAO, we assessed the intracellular accumulation of aminoacetone for the wild-type strain, *aaoSo-mutT*, and *aaoSo* mutants. However, cytoplasmic aminoacetone was not detected in any strain, probably due to the instability of this compound. Using a method described by Kazachkov and Yu, we generated a stable derivative of aminoacetone by addition of 9-fluorenylmethyl chloroformate (FMOC-Cl) to the cellular lysate [23]. As shown in Fig. S5, levels of aminoacetone as much as double that observed in wild-type (76.8±12.5 pmol mg^{-1} cell) were determined in *aaoSo-mutT* mutant (159.8±9.2 pmol mg^{-1} cell) and *aaoSo* strain (147.2±21.4 pmol mg^{-1} cell). Similarly elevated levels were observed in semicarbazide-sensitive amine oxidase (SSAO)-repressed human tissues [23]. This determines that LAAO has a function similar to that of amine oxidase in removing the endogenous aminoacetone.

Next, we added 5 mmol aminoacetone to cultures of wild-type, *aaoSo-mutT* mutant, and *aaoSo* mutant. It was found that aminoacetone reduced the growth of *aaoSo-mutT* mutant by 11%, but it increased the growth of the wild-type and *aaoSo* mutant by 10% (Table 3). Cellular aminoacetone can be derived from threonine or glycine [24], and we supplied threonine to the mutant cultures. Effects of threonine on growth were similar to those of aminoacetone (data not shown). To confirm that the accumulation of threonine-derived aminoacetone reduces the growth of *S. oligofermentans*, we first constructed a threonine dehydrogenase gene (*adh*) deletion mutant. Under *Adh* background conditions, we readily obtained an *aaoSo-mutT* double gene mutant; however, no such mutant was ever made from the wild-type strain as mentioned above. Again this demonstrates that threonine-derived aminoacetone is the deleterious compound that needs to be removed. In addition, the triple mutant *Adh-AaaoSo-MutT* exhibited a growth reduction of only 10% relative to the wild-type strain (Fig. S6). This suggests that the partial growth impairment of *aaoSo-mutT* strain is caused by the accumulation of aminoacetone.

**Discussion**

In this study, we used genetic and biochemical approaches to demonstrate that the gene operon *aaoSo-mutT* constitutes a dual safeguard mechanism, protecting cells from ROS stress in *S. oligofermentans*. Using *aaoSo*-encoded aminoacetone oxidase, cells can reduce the generation of aminoacetone-derived superoxide anion, and the coexpressed MutT eliminates the mutagens derived from ROS. Concerted action of the two proteins can be modulated by the equivalent expression of two genes under the control of a single promoter. This anti-ROS defense mechanism may also be employed by other streptococci. A similar gene operon has also been found in *Streptococcus pneumoniae*, a streptococcus inhabiting in human upper respiratory tract [25]. However, this operon has not yet been found in other lactic acid bacteria whose genomes are available.

### Table 3. Effect of aminoacetone on the growth of *S. oligofermentans* wild-type and two mutants.

| Growth mass (OD600) * | growth time | wild-type | *aaoSo* | *aaoSo-mutT* |
|-----------------------|-------------|-----------|---------|--------------|
|                       | 6 h         |           |         |              |
|                       | 8 h         |           |         |              |

*Symbol: +, 5 mM aminoacetone added; -, no aminoacetone added; Data are shown as means ±SD from three independent experiments.

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Previous research has shown that aminoacetone, a prooxidant, rapidly reacts with oxygen through a superoxide-propagated mechanism even in the presence of strong chelators [26], while semicarbazide-sensitive amine oxidase (SSAO) functions as a cleaner of aminoacetone in human cells [15]. Considering that (1) both LAAO and SSAO belong to the family of amine oxidases; (2) amino acids can also be classified as amines; and (3) aminoacetone is an analog of amino acids; we tested the aminoacetone oxidase activity of LAAO. Pronounced aminoacetone oxidase activity was detected for LAAO, although it was first identified as an amino acid oxidase [11]. Cellular aminoacetone can be derived either from threonine or a condensation of glycine and acetyl-CoA [24,27]. Threonine dehydrogenase, a key enzyme in L-threonine metabolism, catalyzes the conversion of L-threonine to 2-amino-3-ketobutyrate, and the latter is spontaneously decarboxylated to aminoacetone [28]. No gene homology of 2-amino-3-ketobutyrate coenzyme A ligase, which catalyzes glycine and acetyl-CoA in the formation of 2-amino-3-ketobutyrate, was found in the genome of S. oligofermentans. However, the tdh gene was present and an elevated transcription level was detected in both aaoSo-mutT and aaoSo-mutT double gene mutant was obtained using a background of tdh mutation, indicating that aminoacetone is derived from threonine can accumulate to lethal levels when both aaoSo and mutT are absent. Because aminoacetone can react with transition metals, it does not accumulate to a great extent in the cells. This is also true of human cells [23]. The reason why aaoSo inactivation induces expression of tdh remains unknown.

It is known that ROS can oxidize nucleotides [19,29]. Among nucleotides, dGTP is most susceptible to oxidation, especially to hydroxyl radicals, and the oxidized product 8-oxo-dGTP causes mutagenesis and carcinogenesis [18,19]. Organisms have evolved many different approaches to dealing with mutagenesis. These include the MutT protein in E. coli and hMTH-1 protein in humans, both of which are known to hydrolyze 8-oxo-dGTP to the non-harmful 8-oxo-dGMP and PPI [19,30]. MutT belongs to the Nudix hydrolase superfamily, which requires Mg2+ [31]. These enzymes hydrolyze the nucleoside diphosphate modified with other moieties. The most suitable substrate for E. coli MutT is 8-oxo-dGTP [31]. In the present study, similar types of activity were determined for S. oligofermentans MutT. The high levels of expression of mutT in living S. oligofermentans cells and the enzymatic activity of MutT suggest that S. oligofermentans MutT also plays a role in the elimination of deleterious oxidized nucleotides.

Although the aaoSo mutant generates hydroxyl radicals from superoxide anion and H2O2, the intact mutT may be sufficient to the scavenging nucleotide oxides generated by hydroxyl radicals. This may be why no impairment of growth was observed. Under normal growth conditions, the production of ROS is limited to a very low level, thus, AamutT did not show any detectable reduction in growth. However, inactivation of both aaoSo and mutT is lethal for S. oligofermentans. Therefore, synergy of LAAO and MutT is essential for bacterial defense against ROS damage. Here, we propose a working model of LAAO and MutT in prevention of ROS-generated cell oxidative damage (Fig. 4).

The aaoSo-mutT operon and possible coexpression are found in the genome of S. pneumoniae (GenBank accession No. NC_000535). This clinical streptococcus resembles S. oligofermentans with respect to H2O2 production and tolerance [10,25]. Like S. pneumoniae, which lives in an oxygen-rich niche in the upper respiratory tract, S. oligofermentans inhabits the human oral cavity, where the oxygen content fluctuates [25,32,33]. In this way, the aaoSo-mutT operon may have been the evolutionarily selected by bacteria living in environments rich in oxygen.

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**Figure 4. Depicted concerted working model of LAAO and MutT in preventing ROS-generated oxidative damage to cells.** aaoSo and mutT constitute an operon in S. oligofermentans and are co-transcribed, creating the equivalent protein products of the two. The absence of LAAO, an aminoacetone oxidase, was found to cause the accumulation of aminoacetone, which can be derived from threonine by threonine dehydrogenase (Tdh). It generates superoxide anions in the presence of oxygen and transition metals. Fe3+ released from proteins containing [4Fe–4S]2+ cluster by superoxide anion attack was found to trigger the Fenton reaction to form hydroxyl radicals from H2O2, which is produced in abundance from lactate and pyruvate by Lox and Pox, respectively. These hydroxyl radicals oxidize nucleic acids, like dGTP to 8-oxo-dGTP, a mutagen that can cause severe gene mutations. The coexpressed MutT hydrolyzes 8-oxo-dGTP to the harmless 8-oxo-dGMP, and so prevents cell damage. LAAO, L-amino acid oxidase; MutT, pyrophosphohydrolase; Tdh, threonine dehydrogenase; Lox, lactate oxidase; Pox, pyruvate oxidase; Mn-SOD, manganese-dependent superoxide dismutase. Thick arrows refer to the main flux of H2O2; broken arrows indicate the conditional pathways.

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Materials and Methods

Bacterial strains and Culture Conditions

*S. oligofermentans* strain AS 1.3089³ and the *aaoSo*-insertional mutant were cultured in our laboratory [9,11]. Strains were routinely cultured in TYG broth at 37°C in static culture [34]. Agar plates were cultured in a candle jar. Brain heart infusion (BHI) medium (Becton Dickinson, U.S.) supplemented with spectinomycin (800 µg ml⁻¹) or kanamycin (500 µg ml⁻¹) was used for selection of antibiotic-resistant colonies. *E. coli DH5α* (Transgen Biotech, Beijing, China) and BL21 (Novagen, Germany) were grown in Luria–Bertani (LB) broth or on LB agar plate aerobically at 37°C. They were used for plasmid amplification and protein overexpression. Transforms were selected in LB medium with ampicillin (100 µg ml⁻¹), kanamycin (50 µg ml⁻¹), or spectinomycin (250 µg ml⁻¹).

Construction of *aaoSo* mutT, and sodA Gene Deletion Mutants

The genome DNA of *S. oligofermentans* was extracted and purified as described previously [33,35]. PCR primers listed in Table 4 were designed according to the genome draft of *S. oligofermentans* and synthesized by Sangon Company (Shanghai, China). The deletion mutants of the *aaoSo mutT*, and *sodA* genes were constructed using a PCR ligation method [36]. Two ~600 bp sequences upstream and downstream of each of the three genes were PCR amplified using the primers listed in Table 4. PCR products were digested with BamHI (New England Biolabs, U.S.) and a spectinomycin resistance gene cassette was cut from plasmid pDH124 with BamHI as well [36]. The three fragments at equivalent molar ratios were ligated using T₄ DNA ligase (Fermentas, Canada). The fused fragment was transformed into wild-type *S. oligofermentans* with the assistance of a chemical synthetic *S. oligofermentans*-specific competence-stimulating peptide (CSP) [33]. Briefly, overnight culture of *S. oligofermentans* was diluted 1:30 into fresh BHI broth (with 0.4% BSA) and cultured at 37°C for 30 min. The fused fragment was added at a final concentration of 500 µg ml⁻¹ together with the CSP (final concentration 160 ng ml⁻¹). After 5 h incubation, the mixture was plated on BHI agar containing the selective antibiotics and incubated at 37°C in a candle jar. A sodA deletion of *AaaoSo-mutT* mutant was constructed using the same method. Positive transforms of the *AaaoSo-mutT-AaodA* mutant were selected on BHI agar plates containing 500 µg ml⁻¹ kanamycin and 800 µg ml⁻¹ spectinomycin. Transforms of *AaaoSo, AaodA*, and *AaodA* were selected with 500 µg ml⁻¹ each of kanamycin. They were all confirmed by PCR.

Construction of *aaoSo-mutT* and *tdh-aaoSo-mutT* Deletion Mutants

To delete the *aaoSo*-mutT genes on the chromosome, we first constructed a shuttle plasmid, pDL278-*aaoSo-mutT*. The entire *aaoSo-mutT* operon fragment (including the promoter region) was amplified with paired primers pDL-lm-F and pDL-lm-R (Table 4). The fragment was then digested with BamHI and SalI and inserted into plasmid pDL278, which was digested with the same enzymes. The recombinant plasmid was transformed into wild-type strain of *S. oligofermentans*, and the positive strain was selected on BHI agar containing spectinomycin (800 µg ml⁻¹). Upon PCR confirmation of the introduced plasmid-carryed *aaoSo-mutT* operon, a deletion of *aaoSo-mutT* operon on the chromosome was conducted using the same method as the single gene mutation above except that the positive transforms were selected on BHI agar plates containing 500 µg ml⁻¹ kanamycin and 800 µg ml⁻¹ spectinomycin. To produce the triple-gene *tdh-aaoSo-mutT* deletion mutant, the *tdh* gene was knocked out to stop the generation of aminooacetone. Briefly, two ~600 bp sequences upstream and downstream of the *tdh* gene were PCR amplified using the corresponding primers listed in Table 4. PCR products were double digested with *HindIII* and *SmaI* (New England Biolabs, U.S.) and a spectinomycin resistance gene cassette was cut from plasmid pFW3-loc with *HindIII* and *SmaI* [33,37]. The three fragments at equivalent molar ratio were ligated using *T4* DNA ligase (Fermentas, Canada), and the fused fragment was transformed into wild-type *S. oligofermentans*. *Aadh* mutant was selected on BHI agar plate containing 800 µg ml⁻¹ spectinomycin. Under *Aadh* background, the *aaoSo-mutT* operon was knocked out. The triple gene deletion *Aadh-AaaoSo-mutT* was selected on BHI agar plate containing 500 µg ml⁻¹ kanamycin and 800 µg ml⁻¹ spectinomycin. The positive clones were confirmed by PCR.

Construction of *aaoSo-mutT* Gene Complementary Strain

The *aaoSo-mutT* coding region and its inherent promoter were amplified using the paired primers pDL-lm-F and pDL-lm-R (Table 4). The PCR product was double digested with *BamHI* and *SalI* and inserted into plasmid pDL278, which was digested with the same enzymes. Recombinant plasmid pDL278-*aaoSo-mutT* was transformed into the *AaaoSo-mutT* mutant; transforms were selected on BHI agar containing spectinomycin (5 mg ml⁻¹) under aerobic conditions. Positive transforms were confirmed by PCR and sequencing.

Quantification of Gene Expression with Quantitative PCR

Total RNA was extracted using TRIZol reagent (Invitrogen, U.S.) according to the supplier’s recommendations. Cells were harvested at the middle logarithmic phase, washed with RNase-free water, and frozen in liquid nitrogen, and then ground three times with a glass rod. Then TRIZol reagent was added. After incubation at room temperature for 5 min, chloroform was mixed with the lysate and the mixture centrifuged at 12,000 xg. RNA in the aqueous phase was then precipitated using isopropyl alcohol, washed with 75% ethanol, and air dried for 15 min. The pellet was then dissolved in 30 µl RNase-free water, and trace amounts of DNA were removed by DNase. The concentration and quality of RNA samples were determined using NanoDrop spectrophotometer and gel electrophoresis. cDNA was synthesized from 2 µg RNA according to manufacturer’s instructions (Promega, U.S.) and used for quantitative PCR amplification with the corresponding primers listed in Table 4. Amplifications were performed with ABI PRISM 7000 Sequence Detection System (Applied Biosystems, U.S.). The thermocycling parameters included one cycle of 95°C for 10 s, followed by 40 cycles of 95°C for 5 s, 51°C for 30 s, and 72°C for 30 s. Transcripts were quantified with the comparative threshold cycle (Ct) values. To estimate the copy numbers for a given mRNA, a standard curve of the tested gene was generated by cloning the corresponding PCR fragment at a length of about 200 bp into pMD-18T vector (TaKaRa, Japan). The plasmid was then serially diluted and used to generate the standard curve of quantitative PCR. The 16S rRNA gene was used as the biomass reference. The copy number of each gene was normalized to 16S rRNA copies. The copies of the transcript of each gene are shown as per 10,000 16S rRNA copies.

Overexpression and Purification of MutT Protein

The *mutT* gene of *S. oligofermentans* was amplified using a pair of primers (Table 4). Pyrobest DNA polymerase (TaKaRa, Japan) was used for the PCR amplification with the corresponding primers listed in Table 4. The PCR product was double digested with *HindIII* and *SmaI* (New England Biolabs, U.S.) and a spectinomycin resistance gene cassette was cut from plasmid pFW3-loc with *HindIII* and *SmaI* [33,37]. The three fragments at equivalent molar ratio were ligated using *T4* DNA ligase (Fermentas, Canada), and the fused fragment was transformed into wild-type *S. oligofermentans*. *Aadh* mutant was selected on BHI agar plate containing 800 µg ml⁻¹ spectinomycin. Under *Aadh* background, the *aaoSo-mutT* operon was knocked out. The triple gene deletion *Aadh-AaaoSo-mutT* was selected on BHI agar plate containing 500 µg ml⁻¹ kanamycin and 800 µg ml⁻¹ spectinomycin. The positive clones were confirmed by PCR.

The genome DNA of *S. oligofermentans* was extracted and purified as described previously [33,35]. PCR primers listed in Table 4 were designed according to the genome draft of *S. oligofermentans* and synthesized by Sangon Company (Shanghai, China). The deletion mutants of the *aaoSo mutT*, and *sodA* genes were constructed using a PCR ligation method [36]. Two ~600 bp sequences upstream and downstream of each of the three genes were PCR amplified using the primers listed in Table 4. PCR products were digested with *BamHI* (New England Biolabs, U.S.) and a spectinomycin resistance gene cassette was cut from plasmid pDH124 with *BamHI* as well [36]. The three fragments at equivalent molar ratios were ligated using *T4* DNA ligase (Fermentas, Canada). The fused fragment was transformed into wild-type *S. oligofermentans* with the assistance of a chemical synthetic *S. oligofermentans*-specific competence-stimulating peptide (CSP) [33]. Briefly, overnight culture of *S. oligofermentans* was diluted 1:30 into fresh BHI broth (with 0.4% BSA) and cultured at 37°C for 30 min. The fused fragment was added at a final concentration of 500 µg ml⁻¹ together with the CSP (final concentration 160 ng ml⁻¹). After 5 h incubation, the mixture was plated on BHI agar containing the selective antibiotics and incubated at 37°C in a candle jar. A sodA deletion of *AaaoSo-mutT* mutant was constructed using the same method. Positive transforms of the *AaaoSo-mutT-AaodA* mutant were selected on BHI agar plates containing 500 µg ml⁻¹ kanamycin and 800 µg ml⁻¹ spectinomycin. Transforms of *AaaoSo, AaodA*, and *AaodA* were selected with 500 µg ml⁻¹ each of kanamycin. They were all confirmed by PCR.
was used for PCR amplification of *mutT*. A 453 bp fragment was subsequently digested with *Nco*I-*Xho*I (New England Biolabs, U.S.) and cloned into the *Nco*I-*Xho*I restriction sites of the expression vector pET-28a (Novagen, Germany). The His6-tag fusion sites and nucleotide sequences of the double-stranded template DNA were confirmed by sequencing.

The pET-28a plasmid carrying the *mutT* gene was transformed into *E. coli* BL21 (DE3) pLysS (Novagen, Germany) and cultured in LB medium supplemented with 50 μg ml⁻¹ kanamycin. Cells were grown at 37°C to O₆₀₀ 0.4–0.6. Overexpression of MutT protein was induced by addition of 1 mM isopropyl-β-D- thiogalactopyranoside. The culture was incubated for an additional 3 h before harvesting. Cells were collected by centrifugation at 10,000 ×g for 10 min at 4°C, resuspended in the binding buffer (20 mM sodium phosphate, 0.5 M NaCl, 40 mM imidazole, pH 7.4), and was sonicated for 20 min. After the cell lysate was spun down at 10,000 ×g for 10 min at 4°C, the supernatant was filtered through a 0.22 μm polyvinylidene difluoride membrane (Millipore, U.S.) and then applied to a Histrap column (GE Healthcare, USA) previously equilibrated with the binding buffer. Proteins were eluted by elution buffer (20 mM sodium phosphate, 0.5 M NaCl, 500 mM imidazole, pH 7.4), and fractions of the elution were subjected to 12% SDS-PAGE. The desired protein fractions were desalted by loading onto HiPrep™ 26/10 Desalting (GE Healthcare, U.S.) column and eluted with 20 mM Tris–HCl buffer (pH 9.0) with 10% glycerol. The purified MutT protein was stored at –80°C.

### Table 4. Primers used in this study.

| Primer          | Sequence*         | Target genes                  |
|-----------------|-------------------|-------------------------------|
| spacer-lm-F     | TCTCCACCTTGGCGGTTGA | spacer between *aaoSo* and *mutT* |
| spacer-lm-R     | CATGCCACAGTTGACAAATTC |                                    |
| spacer-neg-F    | TCATCAATATTGGGCAAGC | negative control for space check |
| spacer-neg-R    | ACAAACATTTAGCAGGCTACAC |                                    |
| qPCR-16S-F      | GCCGGTGCTAATAACA   | qPCR for 16S rRNA             |
| qPCR-16S-R      | AGACTCCGGTCCTATT   | qPCR for 16S rRNA             |
| qPCR-Sod-F      | ATGGCAATTTTACCAGATCTAC | qPCR for sodA                  |
| qPCR-Sod-R      | TTCCTGGAGTACTCCCAGGA |                                    |
| qPCR-mutT-F     | GAATTGTCACATGTGCA | qPCR for *mutT*               |
| qPCR-mutT-R     | CGGTTGCAAGTTCATGTA | qPCR for *mutT*               |
| qPCR-mutX-F     | ATTCGTAATAATCAACGTGCG | qPCR for *mutX*               |
| qPCR-mutX-R     | TTCCTGAGTTACTCCCGA | qPCR for *mutX*               |
| mutT-kana-up-F  | GAAGAAAAAGGAGAAAGAAG | upstream fragment of *mutT* for deletion |
| mutT-kana-up-R  | AAGGATCCAAATGAGATGTTGACAG | downstream fragment of *mutT* for deletion |
| mutT-kana-down-F| AAGGATCCCTGTTCCTCCCTTTAAAT | downstream fragment of *mutT* for deletion |
| aaoSo-*,kana-up-F| TGCGAAGATCCTTTTGGTTCAGA | upstream fragment of *aaoSo*-*mutT* for deletion |
| aaoSo-*,kana-up-R| TTGGATCGTTCTGATGTTGACAAAC | downstream fragment of *aaoSo*-*mutT* for deletion |
| aaoSo-*,kana-down-F| TTGGATCACGGCCCTCAACAGGAA | downstream fragment of *aaoSo*-*mutT* for deletion |
| aaoSo-*,kana-down-R| TTATCATATAATGCAAACTTCT | downstream fragment of *aaoSo*-*mutT* for deletion |
| pDL-lm-F        | ATGGATCTTAAAAATGAGCTTAATC | construction of shuttle plasmid |
| pDL-lm-R        | TAGTCCGCATTTTTTCTCAAACCAATAAA |                                    |
| lm-kana-up-F    | GAAGAAAAAGGGAGAAAGAAG | upstream fragment of *aaoSo*-*mutT* for deletion |
| lm-kana-up-R    | AAGGATCCAAAATGAGATGTTGACAG | downstream fragment of *aaoSo*-*mutT* for deletion |
| lm-kana-down-F  | AAGGATCTCAGATTTCTCCCTGGT | downstream fragment of *aaoSo*-*mutT* for deletion |
| lm-kana-down-R  | CAGAGGTCGAAGCGCCCTTG | downstream fragment of *aaoSo*-*mutT* for deletion |
| tdh-add9-up-F   | GGATGCCGACTGGCACATGCAATGC | upstream fragment of tdh for deletion |
| tdh-add9-up-R   | TTACCTTCCTTCCTCCCTCCAAATGAG | downstream fragment of tdh for deletion |
| sod-kana-up-F   | GGTGTTGCAATGCTGAGTCGT | upstream fragment of sodA for deletion |
| sod-kana-up-R   | TTCCCGGCTGTCTAAAATTTTCTGCAG | downstream fragment of sodA for deletion |
| sod-kana-down-F| TTGGATCACGGCCCTCAACAGGAA | downstream fragment of sodA for deletion |
| sod-kana-down-R| CAATATTGTGTATGACCCAGCTG | downstream fragment of sodA for deletion |
| pET-mutT-F      | TTCCATGGGGATGAAACAGACGGAAACC | overexpression of MutT |
| pET-mutT-R      | TTCCGAGTTCTCAAGCTAAAAGC |                                    |

*Underlined sequences indicate the restriction enzyme sites. All primers are original in this study.*

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Assay of Aminoacetone Oxidase Activity for LAAO

The aminoacetone activity of LAAO was determined by measuring H_2O_2 production via horseradish peroxidase (HRP)-coupled assay [11]. The assay mixture (1 ml) contained 200 mM LAAO and 25 mM aminoacetone (Toronto Research Chemicals, Canada) in 20 mM Tris–HCl buffer (pH 8.0). Lysine at concentrations of 25 mM and 100 mM was used as the control. The reaction mixture was incubated at 37°C for 30 min. H_2O_2 was quantified using a modified version of a previously described method [11,30]. Briefly, 650 μl of reaction supernatant was added to 600 μl of the solution containing 2.5 mM 4-amino-antipyrine (4-amino-2, 3-dimethyl-1-phenyl-3-pyrazolin-5-one; Sigma–Aldrich, U.S.) and 0.17 M phenol. The reaction proceeded for 4 min at room temperature, and HRP (Sigma–Aldrich, U.S.) was added at a final concentration of 50 nM U ml^-1 in 200 mM potassium phosphate buffer (pH 7.2). After 4 min of incubation at room temperature, OD_325 was measured using a Unico 2100 visible spectrophotometer (Shanghai, China). A standard curve was generated with H_2O_2.

Mn-SOD Activity Assay

The Mn-SOD activity of _S. oligofermentans_ was determined using the method of inhibition of pyrogallol auto-oxidation [39,40]. The reaction mixtures consisted of 4.5 ml 50 mM Tris–HCl (pH 8.2), 1 mM EDTA, 0.1 mM pyrogallol, and 10 μl freshly prepared lysate supernatant of the strains. The same volume of 10 mM HCl was used as a blank control. After mixing, OD_325 was read every 0.5 min at 25°C using a Unico 2100 visible spectrophotometer. The OD_325 increase rate was calculated from 0 to 4 min. One unit of enzymatic activity was defined as the amount of enzyme used for 50% reduction of pyrogallol auto-oxidation rate per min per mg protein.

_S. oligofermentans_ MutT Activity Assay

The enzymatic activity of _S. oligofermentans_ MutT was determined based on the 8-oxo-dGTP degradation [39,40]. The assay mixture contained 100 μl 100 mM Tris–HCl (pH 9.0), 5 mM MgCl_2, 100 μM 8-oxo-dGTP (eENZYME, U.S.), 5% glycerol and 100 nM MutT protein. Desalting buffer without enzyme was used as a negative control. The reaction mixtures were incubated at 37°C for 20 min, and terminated by adding 50 mM EDTA (EDTA: Mg^{2+} = 3.3: 1). Twenty microliters of sample was analyzed on HPLC with the following parameters, flow phases: 20 mM sodium phosphate buffer (pH 6.0) and acetonitrile (95:5); column temperature 30°C; flow rate 1 ml min^-1. The UV absorbance of nucleotide was detected at 293 nm.

Determination of Cellular Aminoacetone Derivate

Cells of _S. oligofermentans_ wild-type and two mutants at their mid-log phase were collected and washed twice with 100 mM sodium phosphate buffer (pH 7.5) by centrifugation at 12,000 x g for 10 min. Fifty milligrams of pelleted cells of each culture were resuspended in 1 ml 100 mM phosphate buffer (pH 7.3), then homogenized in a MINI Bead Beater (Biospec Products; Bartlesville, OK, U.S.) for 20 s, with a 2-min interval on ice. This process was repeated five times. After centrifugation of the cell extract at 12,000 x g for 10 min at 4°C, the supernatant was partially purified using a 3 KD ultrafiltration tube (MWCO, Millipore, U.S.). Synthetic aminoacetone (Toronto Research Chemicals, Canada) and the cell extract samples were treated with FMOC-Cl to obtain the aminoacetone derivates using a modified version of the method described by Kazachkov and Yu [23]. Briefly, 1 ml sodium borate buffer (0.4 M, pH 10) was added to 1 ml of samples or aminoacetone (10 mM) and vortexed for 1 min. Then, 1 ml FMOC-Cl reagent (10 mM in acetonitrile) was mixed with the samples by vortex for 1 min. Five milliliters of hexane was added to terminate the reaction by extracting the excess reagent (FMOC-Cl), hydrolysis product FMOC-OH and acetonitrile. The upper hexane layer was discarded and the process was repeated twice, and the remaining solution was neutralized by addition of 100 μl 20% (v/v) acetic acid for HPLC assay using a modified version of the method described by Xiao and Yu [41]. The assay was performed on Shimadzu HPLC system (Japan) with a C18 reverse phase column (4.6 x 250 mm/ 5 μm; Agilent, U.S.), and the compound separation was achieved under the following conditions, flow phases: 5 μM potassium phosphate buffer (pH 4.40–4.45) and acetonitrile (60:40); column temperature 30°C; flow rate 1 ml min^-1. FMOC-aminoacetone was detected for UV absorbance at 265 nm.

Determination of ROS Levels

Cellular ROS levels in _S. oligofermentans_ were measured using a spin trapping agent combined with ESR spectroscopy (Bruker ESP 300, Germany) [21,22]. 5,5-Dimethyl-l-pyrroline N-oxide (DMPO, TCI, Japan) was purified by stirring a solution (0.25 M) of the spin trap in deionized water with a little activated charcoal for 1 hour. After filtration, this solution was stored in the dark at -20°C. The wild-type strain and two mutants were collected at mid-log phase and washed twice with sodium sulfate solution (50 mM). The standard incubation consisted of the following final concentrations: 30 mg bacterial cells, DMPO solution (100 mM), and each sample were brought up to 0.5 ml with deionized water. The mixtures were incubated at 37°C for 5 min before detection. ESR samples in a quartz flat cell were fixed in the cavity of the ESR spectrometer. ESR measurements were taken at 9.75 GHz with a 100 kHz modulation frequency, and the following ESR spin trapping parameters were used: field sweep, 343.2–353.2 mT; microwave frequency, 9.75 GHz; sweep time, 168 s; microwave power, 12.8 mW; modulation amplitude 1 mT; and time constant, 328 ms.

Supporting Information

**Figure S1** Growth of the wild-type, _AaaoSo_ and _AaamutT_ mutants of _S. oligofermentans_ in glucose. Strains were pre-cultured overnight in TYG medium, and 1:100 diluted with fresh TYG medium with the same OD_600_. Samples were taken every 1 h to detect OD_600_. Symbols: Δ, wild-type strain; ○, _AaaoSo_ mutant; ×, _AaamutT_ mutant. The results are shown as the means±SD of three independent experiments. (TIF)

**Figure S2** Enzymatic assays for the overexpressed LAAO of _S. oligofermentans_. Activities are expressed by H_2O_2 production rate as described in materials and methods. Data are the means of three assays, and standard deviations are shown. (TIF)

**Figure S3** Pyrophosphohydrolase activity of _S. oligofermentans_ MutT for 8-oxo-dGTP determined through HPLC. (A) 8-oxo-dGTP (100 μM) and (B) 8-oxo-dGTP incubated with 100 nM purified MutT at 37°C for 20 min. Retention times for 8-oxo-dGTP and 8-oxo-dGMP are shown. (TIF)

**Figure S4** Growth of the wild-type, _AaaoSo-mutT, Asod_, and _AaaoSo-mutT-Asod_ mutants of _S. oligofermentans_. Strains were cultured overnight and adjusted to the same OD_600_, and then 1:100 diluted with fresh TYG medium and cultured.
statically unless indicated otherwise. OD_{600} was measured in 1 h interval. Symbols: Δ, wild-type strain; □, AaoSo-mutT mutant; ○, Asod mutant; ×, AaoSo-mutT-Asod mutant; •, AaoSo-mutT-Asod mutant growing strictly anaerobically. Results represent the means ± SD from three independent experiments. (TIF)

Figure S5 HPLC chromatogram of FMOC-amineacetone obtained from the wild-type and both mutants cells of *S. oligofermentans*. The FMOC derivate signal of the chemical amineacetone appears at 27.5 min under the analytical conditions. (A) FMOC derivate of chemical amineacetone; (B) wild-type strain; (C) AaoSo mutant, and (D) AaoSo-mutT mutant. (TIF)

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**Figure S6** Growth of *Attdh-AaoSo-mutT* mutant and the wild-type of *S. oligofermentans*. Strains were cultured overnight and adjusted to the same OD_{600} in TYG medium, and 1:100 diluted with the same medium. OD_{600} was detected in 1 h interval. Symbols: Δ, wild-type strain; •, *Attdh-AaoSo-mutT* mutant. The results are shown as the means ± SD of three independent experiments. (TIF)

**Author Contributions**
Conceived and designed the experiments: PZ XD HT. Performed the experiments: PZ LL. Analyzed the data: PZ XD HT. Contributed reagents/materials/analysis tools: PZ LL. Wrote the paper: PZ XD HT.

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