A Novel 5-Enolpyruvylshikimate-3-Phosphate Synthase from *Rahnella aquatilis* with Significantly Reduced Glyphosate Sensitivity

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**Abstract**

The 5-enolpyruvylshikimate-3-phosphate synthase (EPSPS; EC 2.5.1.19) is a key enzyme in the shikimate pathway for the production of aromatic amino acids and chorismate-derived secondary metabolites in plants, fungi, and microorganisms. It is also the target of the broad-spectrum herbicide glyphosate. Natural glyphosate resistance is generally thought to occur within microorganisms in a strong selective pressure condition. *Rahnella aquatilis* strain GR20, an antagonist against pathogenic agrobacterial strains of grape crown gall, was isolated from the rhizosphere of grape in glyphosate-contaminated vineyards. A novel gene encoding EPSPS was identified from the isolated bacterium by complementation of an *Escherichia coli* auxotrophic aroA mutant. The EPSPS, named AroA<sub>aquatilis</sub>, was expressed and purified from *E. coli*, and key kinetic values were determined. The full-length enzyme exhibited higher tolerance to glyphosate than the *E. coli* EPSPS (AroA<sub>E.coli</sub>), while retaining high affinity for the substrate phosphoenolpyruvate. Transgenic plants of AroA<sub>aquatilis</sub> were also observed to be more resistant to glyphosate at a concentration of 5 mM than that of AroA<sub>E.coli</sub>. To probe the sites contributing to increased tolerance to glyphosate, mutant *R.aquatilis* EPSPS enzymes were produced with the c-strand of subdomain 3 and the f-strand of subdomain 5 (Thr38Lys, Arg40Val, Arg222Gln, Ser224Val, Ile225Val, and Gln226Lys) substituted by the corresponding region of the *E. coli* EPSPS. The mutant enzyme exhibited greater sensitivity to glyphosate than the wild type *R.aquatilis* EPSPS with little change of affinity for its first substrate, shikimate-3-phosphate (S3P) and phosphoenolpyruvate (PEP). The effect of the residues on subdomain 5 on glyphosate resistance was more obvious.

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**Introduction**

The enzyme 5-enolpyruvylshikimate-3-phosphate synthase (EPSPS; EC 2.5.1.19) is a carboxyvinyl transferase that catalyzes the transfer of the enolpyruyl moiety of phosphoenolpyruvate (PEP) to the 5-hydroxyl of shikimate-3-phosphate (S3P) and forms enolpyruvylshikimate-3-phosphate. It is the sixth key enzyme in the shikimate pathway, which is essential for the production of aromatic amino acids Phe, Tyr, and Trp, as well as chorismate-derived secondary metabolites in plants, fungi, and microorganisms [1]. EPSPS has attracted significant attention since it was identified as the primary target of glyphosate in the 1980s [2]. Competitive with PEP, glyphosate forms a stable ternary complex with the enzyme and S3P [3,4]. The safety and efficacy of glyphosate, together with the widely used glyphosate-resistant crop plants containing a transgene for EPSPS, have combined to make glyphosate the most used herbicide in the world. Herbicide-resistant corn (*Zea mays* L.), canola (*Brassica napus* L.), cotton (*Gossypium hirsutum* L.), alfalfa (*Medicago sativa* L.), and soybeans account for about 82% of biotech crops.

In the past years, a number of glyphosate-resistant bacteria and glyphosate-tolerant plants have been identified. EPSPS from different organisms has been divided into two classes according to intrinsic glyphosate sensitivity. Class I EPSPS was generally identified from plants and bacteria, such as *Escherchia coli* and *Salmonella typhimurium*, the catalytic activity of which was inhibited at low micromolar concentrations of glyphosate [5]. However, the tolerance to glyphosate can be generated by different mutations [6,7,8]. For example, alterations of Gly96Ala in *E. coli* and Pro101Ser in *S. typhimurium* have been shown to present resistance to glyphosate. Class II EPSPS, identified from some bacteria, such as *Pseudomonas* strain PG2982, *Agrobacterium tumefaciens* strain CP4, *Streptococcus pneumoniae*, and *Staphylococcus aureus*, was distinguished by its capability to sustain efficient catalysis in the presence of high glyphosate concentrations [9,10]. These two types of EPSPS share less than 30% amino acid sequence identity [11].
In vineyards, glyphosate is widely used to control weeds directly beneath the vines. Soil and rhizosphere microorganisms from vineyards can harbor highly glyphosate-tolerant EPSPS. In this study, we have isolated a glyphosate-tolerant Rahnella aquatilis strain from the rhizosphere of grape in glyphosate-contaminated vineyards. The strain showed a significant inhibition to the pathogen causing crown galls, a worldwide plant disease in grape-growing regions [12,13]. The isolation, cloning, and characterization of the R. aquatilis strain EPSPS, AroAR_aquatilis, are described. Although this EPSPS shares 88% amino acid identity with the EPSPS (P07638) from E. coli, it has more excellent kinetic qualities as it possesses similar binding affinity for the natural substrate PEP but lower affinity sensitivity to glyphosate than the E. coli EPSPS. The protein also confers glyphosate tolerance on transgenic plants in greenhouse. Using site-directed mutagenesis, two regions at subdomains 3 and 5 were identified to contribute to increased tolerance to glyphosate.

Results

Isolation and identification of the glyphosate-tolerant strain

After selection on low-concentration glyphosate and further verification on high-concentration glyphosate, one strain has been isolated from the rhizosphere of grape. This strain, named GR20, grew well on M9 plates in the presence of 200 mM glyphosate. GR20 was identified as R. aquatilis by the biolog identification system. It was small, rod shaped, and Gram negative, and the biochemical test results showed that it was negative for cytochrome oxidase and positive for catalase. The 16S rDNA gene sequence (1,506 bp) confirmed that GR20 was strongly related to R. aquatilis (99.8%).

Cloning and Sequence analysis of the EPSPS gene from R. aquatilis GR20

To isolate the gene involved in glyphosate tolerance from R. aquatilis GR20, a genomic library was constructed from this strain. There were about 2×10^5 ampicillin-resistant clones on the LB plates. Selection on M9 plates containing 60 mM glyphosate led to the identification of five clones. The plasmid of all these clones was purified. The translated protein of ORF427 was determined to be 36.7 kDa. Further examine the premise that the AroAR_aquatilis is better in terms of glyphosate resistance than AroAR_E.coli. The plasmid pGR20-5 with about 2.1 kb insert fragment was selected to be sequenced subsequently.

The 2.1 kb insert fragment in pGR20-5 was sequenced and analyzed. A complete open reading frame of 1284 bp with a G + C content of 51.4% was deduced and named AroAR_aquatilis (GenBank sequence accession number GQ499276). To determine whether ORF427 was translated, AroAR_aquatilis was expressed and purified. The translated protein of ORF427 was determined to be 47 kDa by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (PAGE), which agreed well with the molecular size of the predicted product (Fig. S1). The N-terminal amino acid sequence of AroAR_aquatilis was further determined as M-E-S-L-I-I-T-L-Q-P-F-V-A by automated Edman degradation. The result confirmed that the initiator codon of AroAR_aquatilis is GUG.

To determine whether ORF427 is sufficient for glyphosate tolerance, a PCR experiment was carried out to amplify only the sequence encoding ORF427 from pGR20-5. The amplified fragment was cloned into the expression vector pYXP251 and transformed into the E. coli araoA mutant strain ER2799. The transformants were capable of complementing its araoA mutation for growth on M9 minimal medium supplemented with 60 mM glyphosate (Fig. 1).

Nucleotide sequence analysis indicated that AroAR_aquatilis shared more than 80% amino acid identity with E. coli EPSPS (AroAR_E.coli; Fig. 2A). Detected from the structure of AroAR_aquatilis modeled by SWISS-MODEL based on the structures of AroA E. coli (PDB 1G6S), the most different amino acids between AroAR_aquatilis and AroAR_E.coli were lined on four-stranded β-sheet structures (159–165; 194–193; 213–218; 221–226) in subdomain 5 [14] and the c-strand of subdomain 3 (Fig. 2B). Phylogenetic tree analysis indicated that AroAR_aquatilis belonged to class I EPSPS (Fig. 2C).

Growth of cells in the presence of glyphosate

The growth curves of the E. coli araoA-deleted mutant ER2799 harboring plasmid p251-AroARa or p251-AroAEc are shown in Fig. 3. Cells were grown in liquid M9 minimal medium containing various concentrations of glyphosate. The results show that the growth of cells harboring p251-AroAEc was strongly inhibited by 50 mM glyphosate. In contrast, cells harboring p251-AroARa can grow at 150 mM glyphosate. The result suggests that AroAR_aquatilis was more highly tolerant to glyphosate than AroAR_E.coli. May be at least one sequence domain at the different regions of subdomain 5 and subdomain 3 is related with glyphosate resistance activity. To determine why AroAR_aquatilis had a higher glyphosate tolerance than AroAR_E.coli, the domains of c-strand of subdomain 3, which is near the hinge region for the relationship of the EPSPS two globular structure, and the f-strand of subdomain 5, which was induced to chemical shift changes upon S3P binding, were selected to examine. The residues in the two domain of AroAR_aquatilis were changed into the corresponding amino acids of AroAR_E.coli by multiple-site mutagenesis method (Thr38Lys and Arg40Val, Arg222Gln, Ser224Val, Ile225Val, and Gln226Lys). Cells expressing the regions mutant m35 AroAR_aquatilis grew much worse than the cells expressing the wild type. The growth potential of the cells expressing m35 AroAR_aquatilis was similar to that of cells expressing AroAR_E.coli in the presence of 100–150 mM glyphosate (Fig. 3).

Assay for glyphosate tolerance of transgenic tobacco

Further examine the premise that the AroAR_aquatilis is better in terms of glyphosate resistance than AroAR_E.coli was confirmed in transgenic tobacco. Expression cassettes comprising a CaMV35S promoter, chloroplast transit sequence of tobacco, and EPSPS coding region were cloned into the T-DNA region of the binary vector pYF7716 [15] (Fig. 4A). Then araoA genes were introduced to tobacco plants via Agrobacterium-mediated transformation. Transgenic plants were obtained from plantlets regenerated on medium containing 30 mg/L hygromycin and further confirmed by PCR analysis. Sixteen AroAR_aquatilis transgenic plants survived after spraying twice with 5 mM glyphosate. Three AroAR_aquatilis transgenic lines (Ra3, Ra8, and Ra11) and two AroAR_E.coli transgenic lines (Ec3 and Ec7) were analyzed for gene expression using RT-PCR analysis. The results of RT-PCR showed that the inserted genes were actively and stably transcribed in transgenic plants (Fig. 4B).

Subsequently, the glyphosate resistance of transgenic plants was compared at different growth stages. The leaf disc of the transgenic line containing AroAR_aquatilis had no visible damage at a concentration of 0.4 mM glyphosate, whereas the leaf disc of the transgenic lines containing AroAR_E.coli was mortified severely on the periphery. Untransformed control plants died completely at a concentration of 0.1 mM (Fig. 5). The transgenic lines of AroAR_aquatilis and AroAR_E.coli were also used to test glyphosate resistance by spraying with 5 mM glyphosate in five- to six-leaf stage. AroAR_aquatilis transgenic plants grew well with normal morphology except for chlorosis of old leaves, whereas AroAR_E.coli
transgenic plants showed severe chlorosis on most leaves, and nontransformed plants showed very rapid chlorosis and bleaching and then wilted and died (Fig. 6). These results indicate that AroAR.aquatilis was more resistant to glyphosate exposure than AroAE.coli, and the characteristic of glyphosate resistance can be heritable in transgenic plants.

To prove both identified regions was critical for glyphosate resistance in class I EPSPS, the mutant, m3AroAR.aquatilis, was also transfer to tobacco. After examined by RT-PCR (Fig. 4B), the transgenic lines of AroAR.aquatilis were further assessed for resistance to glyphosate by spraying with glyphosate. In comparison, all transgenic plants containing m5AroAR.aquatilis have a serious injury as transgenic plants containing AroAE.coli after two spraying (Fig. 6).

**Kinetic properties of AroAR.aquatilis**

Sequence analysis showed that it was significantly different between AroAR.aquatilis and AroAE.coli in the subdomains 3 and 5 of the N-terminal domain. Biological experimentation in transformed E. coli and transgenic tobacco plants that compares activity of the constructed mutant gene m5AroAR.aquatilis with that of the wild-type AroAR.aquatilis and AroAE.coli showed that both regions of the c-strand of subdomain 3 (Thr38Lys and Arg40Val) and the f-strand of the subdomain 5 were key domains for glyphosate resistance activity.

All mutant and wild EPSPS enzymes were overexpressed separately in E. coli, then extracted and purified for activity assay. The $K_i$ [PEP] and $K_i$ [S3P] value of AroAR.aquatilis was similar with that of AroAE.coli. However, the $K_i$ [glyphosate] and IC$50$ [glyphosate] of AroAR.aquatilis were approximately 35-fold and 5-fold higher than the values of AroAE.coli respectively (Table 1). These results exemplify that both enzymes have similar affinity for both substrates PEP and S3P but AroAR.aquatilis had a higher glyphosate tolerance and a lower level of affinity for glyphosate than AroAE.coli.

Biochemical analyses of mutants have also allowed some regions to be identified as essential or nonessential for glyphosate resistance activity. Kinetic characterizations of the wild-type AroAR.aquatilis and its mutants on both regions were detected. The $K_{in}$ [PEP] and $K_{in}$ [S3P] of the mutant m3AroAR.aquatilis were similar to the wild type, whereas the values of $K_i$ [glyphosate] and IC$50$ [glyphosate] decreased apparently respect to the wild type enzyme. The values of $K_i$ [glyphosate] and IC$50$ [glyphosate] of m3AroAR.aquatilis are only 6-fold and 2-fold higher than that of AroAE.coli respectively (Table 1). These results indicated that the c-strand of subdomain 3 and the f-strand of the subdomain 5 were important for glyphosate binding and confered a close relationship with glyphosate resistance. To detect what is contribution for glyphosate resistance, more kinetic analysis of the independent mutant on both regions was done. The $K_{in}$ [PEP] and $K_{in}$ [S3P] of the two mutants (m3AroAR.aquatilis and m5AroAR.aquatilis) have no significant change. But the values of $K_i$ [glyphosate] and IC$50$ [glyphosate] decreased when compared with that of the wild type. The ratios of the wild type enzyme over the mutant m3AroAR.aquatilis were approximately 2.8 and 1.5 for $K_i$ [glyphosate] and IC$50$ [glyphosate] respectively, while the ratios over the mutant m5AroAR.aquatilis were 3.7 and 2.0. It indicated that the f-strand of the subdomain 5 was more important for glyphosate resistance than the c-strand of subdomain 3 (Table 1). Further, in order to testify that the f-strand of the subdomain 5 was important for glyphosate binding, we have generated m5AroAE.coli with the subdomain 5 of AroAE.coli. The kinetic characterizations showed that the values of $K_i$ [glyphosate] and IC$50$ [glyphosate] of m5AroAE.coli are 7-fold and 3-fold higher than that of AroAE.coli. It is obvious that the f-strand of the subdomain 5 was indeed important for glyphosate binding.

**Discussion**

In this study, a novel aroA gene was cloned from an R. aquatilis strain isolated from the rhizosphere of grape in glyphosate-
It was shown that the aroA gene product (AroA<sub>R.aquatilis</sub>) could complement <i>E. coli</i> AroA mutations. Sequence analysis revealed that AroA<sub>R.aquatilis</sub> was most closely related to the EPSPS of <i>E. coli</i> AroA and belonged to class I AroA enzymes (Fig. 2C).

In addition to wastewater, puddle, and human clinical specimens, <i>R. aquatilis</i> can also be isolated from nonpolluted soil. Some strains of <i>R. aquatilis</i> were plant rhizosphere-associated bacteria. <i>R. aquatilis</i> has been reported to be a potential biological control agent for different <i>Agrobacterium</i> strains, which are causal organisms of crown gall disease [16]. Although the genus of <i>Rahnella</i> is widely distributed in water and soil, nobody has noticed that <i>R. aquatilis</i> strain have natural glyphosate tolerance until now. <i>R. aquatilis</i> may have acquired an aroA gene from other soil microbial communities or improved glyphosate tolerance of AroA enzymes through natural selection.

We have examined the glyphosate resistance activity of AroA<sub>R.aquatilis</sub> in transformed <i>E. coli</i> and transgenic tobacco. Transformed <i>E. coli</i> and transgenic plants with AroA<sub>R.aquatilis</sub> showed greater glyphosate resistance as compared to AroA<sub>E.coli</sub>. The AroA<sub>R.aquatilis</sub> transgenic plant line can survive treatment with 5 mM glyphosate. Kinetic analysis of the AroA<sub>R.aquatilis</sub> also indicated that it had lower glyphosate affinities than AroA<sub>E.coli</sub>.

The reaction of EPSPS with S3P and PEP and the inhibition of this reaction by glyphosate have been extensively studied during the past 30 years. Glyphosate is a reversible inhibitor of EPSPS because it can bind adjacent to S3P in the PEP binding site and mimic an intermediate state of enzyme-substrates complex. In

Figure 2. Sequence analysis of the AroA protein from <i>Rahnella aquatilis</i> G20. (A) Protein sequence comparison of both AroA protein from <i>R. aquatilis</i> G20 and <i>E. coli</i>. Amino acids for S3P binding are indicated as asterisks, and amino acids for PEP binding are indicated as squares. The arrow shows the β-sheet in the secondary structure of the subdomain 5. Residues that were mutated in this study were also showed. (B) Views down the axis of the N-terminal domain of the model of AroA<sub>R.aquatilis</sub> (AroA-Ra). Each subdomain is numbered according to Stallings et al (30). The proposed S3P and PEP binding residues are rendered in CPK. Residues in chemical shift upon S3P binding are indicated in black. The mutant regions were displayed using blue color. (C) Phylogenetic analysis of AroA constructed using DNAMAN. The tree suggests that AroA<sub>R.aquatilis</sub> belonged to Class I AroA protein. Class I AroA proteins were obtained from <i>E. coli</i> (P07638), <i>Aeromonas salmonicida</i> (Q03321), <i>Arabidopsis thaliana</i> (P05466), <i>N. tabacum</i> (P25981), <i>Petunia hybrida</i> (P11043), Z. mays (CAA44974), and <i>Bordetella pertussis</i> (P12421). Class II AroA proteins were obtained from <i>Pseudomonas</i> strain PG2982 (P56952), <i>A. tumefaciens</i> strain CP4 (Q9R4E4), <i>Bacillus subtilis</i> (P20691), <i>S. aureus</i> (Q05615), and <i>S. pneumoniae</i> (Q9S400). doi:10.1371/journal.pone.0039579.g002
recent years, a large number of active sites of EPSPS have been studied and identified [17,18,19,20]. The glyphosate binding site is dominated by charged residues Lys22, Arg124, and Lys411, which have previously been concerned in PEP binding [21]. Mutation of these amino acids can result in a significant change in glyphosate tolerance. Target site glyphosate tolerance can also be induced by other specific mutations, including Thr42Met [22], Gly96Ala [23], Thr97Ile [24], Pro101Leu, Pro101Thr, Pro101Ala, and Pro101S-

Figure 3. Growth curves of the E. coli aroA mutant strain ER2799 cells expressing AroAR.aquatilis (AroA-Ra), m35AroAR.aquatilis (m35AroA-Ra), and AroAE.coli (AroA-Ec) in liquid M9 minimal medium supplemented with glyphosate at the concentrations indicated. The results presented are the averages of two sets of experiments done in triplicate. doi:10.1371/journal.pone.0039579.g003

Figure 4. (A) Schematic diagram of the T-DNA region of the binary vectors used in this study. LB, left border; Hyg, hygromycin phosphotransferase gene; D35S, double cauliflower mosaic virus 35S promoter; TSP, tobacco transit peptide; AroA, EPSPS gene; NOS-T, nopaline synthase gene terminator; RB, right border. For the steady transmission of the aroA gene, scaffold attachment region (SAR) was fused upstream of the D35S promoter. (B) Evaluation of the glyphosate resistance of transgenic lines. (A) RT-PCR analysis of the transcripts of the AroAR.aquatilis, m35AroAR.aquatilis, and AroAE.coli genes in transgenic and nontransgenic tobacco lines. Normalized expression of the Actin gene was used as control. WT, nontransgenic tobacco; Ra3, Ra8, and Ra11, AroAR.aquatilis transgenic lines; Ec3 and Ec7, AroAE.coli transgenic lines; mRa1, mRa4, mRa5, m35AroAR.aquatilis transgenic lines. doi:10.1371/journal.pone.0039579.g004
er [8, 18, 25], and Ala183Thr [23], in which residues 97–105 form an internal helix in the N-terminal globular domain and correct with glyphosate closely, and Ala183 is an important residue for EPSPS-glyphosate interaction. The residues important for S3P binding and PEP binding and the domains for glyphosate tolerance were found to be conservative between AroA<sub>R.aquatilis</sub> and AroA<sub>E.coli</sub>.

As compared to the <i>E. coli</i> AroA protein, AroA<sub>R.aquatilis</sub> has two significant different regions in the N-terminal domain. Although both different regions were at the position distant from the active site for S3P binding and PEP binding, they are related with the conformational change of N- and C-terminal globular domain. One region is located in subdomain 5. Residues in the four-stranded β-sheet structures (159–165, 184–193, 213–218, and 221–226) of this domain are changed greatly (Fig. 2A and B).

Some residues in the f-strand of the subdomain 5 will be induced to chemical shift changes upon S3P binding. As revealed in X-ray crystallography studies, S3P triggers the two globular domains to move toward each other in a screw-like movement and cause its transition from the open to the closed state [21]. These different residues may induce structural adjustments in hydrogen bonding networks and result in positional shifts of the residues connected to the substrate [26]. The other is located in the c-strand of subdomain 3 (Fig. 2A and B). In this region, some residues are near the hinge region, which will alter the relationship of the two globular structures of EPSPS. This proposed mechanism was corroborated with the finding that the substitution of bulkier hydrophobic amino acid methionine for Thr42, a residue near the hinge region, affects the screw-like closure by altering the relative positions of the clusters of the active site amino acids that reside on opposite globular domains [7].

To prove that both regions affected the glyphosate affinities of AroA, the c-strand of subdomain 3 and the f-strand of subdomain 5 of AroA<sub>R.aquatilis</sub> were substituted with the corresponding residues in the same region of AroA<sub>E.coli</sub> by multiple-site mutagenesis method. Both domain replacements decreased the <i>K<sub>i</sub></i> [glyphosate] and IC<sub>50</sub> significantly. However, the <i>K<sub>m</sub></i> [PEP] and <i>K<sub>m</sub></i> [S3P] changed trivially. This implied that glyphosate tolerance of AroA was related to the regions containing residues whose chemical shifts change upon S3 binding and the regions containing residues near the hinge domain. Further experiment proved that the f-strand of subdomain 5 contributed more to glyphosate tolerance than the c-strand of subdomain 3 (Table 1, Fig. S2). It suggested that the open conformation of the enzyme-S3P complex will affect the glyphosate affinities.

Although a large number of AroA enzymes have been identified and tested as glyphosate resistant, only AroA variants derived from the <i>A. tumefaciens</i> strain CP4 have been successfully used commercially [10]. Many single-site mutations have been made to investigate the molecular basis of the glyphosate insensitivity of the AroA enzyme. However, glyphosate tolerance is often paralleled by a decrease in the affinity of EPSPS for PEP. Thus, they were insufficient for commercial glyphosate-resistant crops.

Figure 5. Herbicide resistance in leaf discs after 3 weeks in culture with glyphosate. Leaf discs from transgenic plants AroA<sub>R.aquatilis</sub> (a), AroA<sub>E.coli</sub> (b), and nontransformed control plants (c) were cultured on solid MS medium containing 0 to 0.8 mM glyphosate (columns 1–5 contained 0, 0.1, 0.2, 0.4, and 0.8 mM glyphosate, respectively). doi:10.1371/journal.pone.0039579.g005

Figure 6. The transgenic plants were sprayed with glyphosate at a dose of 5 mM. Injury was observed visually 2 weeks after the application. (a) nontransformed plants; (b) transgenic tobacco line Ec3; (c) transgenic tobacco line mRa1; and (d) transgenic tobacco line Ra11. For (B) and (C), the same results were obtained in three independent experiments and are represented by the effects shown here. doi:10.1371/journal.pone.0039579.g006
For example, although the $K_i$ [glyphosate] value of the E. coli EPSPS mutant Gly96Ala was more than 1000-fold higher than that of the wild type, the $K_M$ [PEP] of the mutant increased 23 times [11]. Multisite mutations with more favorable properties were sought and discovered. For example, the double mutant Z. mays EPSPS Thr102Ile/Pro106Ser (abbreviated as TIPS) had particularly favorable characteristics. The TIPS mutations have been introduced into the endogenous EPSPS enzyme of Z. mays to produce the first commercial varieties of glyphosate-resistant maize (U.S. Patent 6,040,497). In this research, we found that AroA from E. coli, had greater glyphosate tolerance without decreasing its affinity for PEP and S3P. AroA from E. coli, may help to obtain a new type of class I AroA proteins that not only show high tolerance to glyphosate but also have high affinity for PEP. This can be done through domain replacement or multi-sites mutation but not single-site mutation. Through site-directed mutation and optimal codon usage, the new aroA gene is likely to improve its efficiency for glyphosate tolerance in engineering crops.

Materials and Methods

Medium and chemicals

S3P, glyphosate (free acid form), and PEP were purchased from Sigma-Aldrich. All other chemicals were of analytical grade. E. coli strain ER2799 (with the EPSPS gene deleted from its genome) was a kind gift from Professor Thomas C. Evans, Jr. (New England Biolabs, USA). Agrobacterium strains were bought from the Agricultural Culture Collection of China.

Study area and ethical approval

The field studies were conducted in a vineyard which located at Hongqi village, Minheng district, Shanghai, China. Glyphosate has been used extensively as a broad-spectrum control agent to kill weed plants over 10 years in this vineyard.

This study involved collection of 10 samples of soil from the rhizosphere of grape in glyphosate used extensively vineyards, without human participants and animal work, therefore the local ethics committee considered that informed verbal consent was appropriate and this was approved by district officials, village, sub-village leaders and the owner of this vineyards. No specific permits were required for the field studies. Also the field studies did not involve endangered or protected species and no specific permissions were required for these locations. And local ethics committee waived the need for consent.

Identification of strain GR20

The morphological, cultural, biochemical, and physiological characteristics of GR20 were examined according to the methods previously reported by Farmer [27]. Chromosomal DNA of GR20 was extracted and purified according to the reported method [28]. The 16S rDNA was amplified from GR20 genomic DNA with the primers 16SR (5'-AGAGTTTGATCTTCGGCTCAG-3') and 16SF (5'-ACGGCTACCTTGTGACGCTT-3'). The PCR fragment was cloned and sequenced.

Isolation of the gene involved in glyphosate tolerance

Chromosomal DNA isolated from strain GR20 was partially digested with Sau3A, and DNA fragments of 2 to 4 kb were purified and inserted into the BamHI site of dephosphorylated pUC18 [29]. A genomic library was constructed by electroporating ligated DNA into E. coli DH5α-compotent cells. Then the genomic library plasmid was transformed to the E. coli araA mutant strain ER2799 and screened on M9 agar plates supplemented with 60 mM glyphosate. One colony harboring a

Table 1. Kinetic properties of AroA from Rahnella aquatilis and E. coli. a

| Enzyme            | Sp actb (U/mg) | $K_m$ [PEP]c (μM) | $K_m$ [S3P]c (μM) | $K_i$ [glyphosate]d (μM) | $IC_{50}$ [glyphosate]e (mM) | $K_i/K_{IC_{50}}$ [PEP]f |
|-------------------|----------------|-------------------|-------------------|-------------------------|----------------------------|-----------------------------|
| AroA_R.aquatilis  | 82.8 ± 3.1     | 78.3 ± 2.3        | 86.3 ± 3.5        | 49.6 ± 5.2              | 0.26 ± 0.02                | 0.63                         |
| m3AroA_R.aquatilis| 66.4 ± 4.4     | 74.5 ± 4.5        | 72.1 ± 6.2        | 17.2 ± 1.9              | 0.17 ± 0.01                | 0.24                         |
| m5AroA_R.aquatilis| 54.3 ± 3.2     | 71.4 ± 3.6        | 69.8 ± 2.1        | 13.5 ± 3.6              | 0.13 ± 0.01                | 0.18                         |
| m35AroA_R.aquatilis| 48.7 ± 4.1    | 66.3 ± 4.2        | 71.2 ± 3.8        | 9.0 ± 0.7               | 0.11 ± 0.01                | 0.14                         |
| AroN_e.coli       | 41.1 ± 5.3     | 65.8 ± 5.6        | 60.1 ± 3.4        | 1.4 ± 0.1               | 0.05 ± 0.01                | 0.021                        |
| m5AroN_e.coli     | 60.3 ± 3.7     | 67.6 ± 2.9        | 65.3 ± 2.9        | 8.4 ± 3.4               | 0.15 ± 0.01                | 0.22                         |
| G96A AroN_e.coli  | 71.4 ± 4.5     | 119.5 ± 5.1       | 61.2 ± 4.5        | 219.3 ± 1.9             | 0.22 ± 0.02                | 0.19                         |
| G96A AroA_R.aquatilis| 100.5 ± 2.9   | 148.3 ± 4.5       | 84.5 ± 5.4        | 450.4 ± 3.4             | 0.34 ± 0.01                | 0.38                         |

aThe results presented are the averages of two sets of experiments done in triplicate.
bSp act, specific activity; 1 unit (U) of EPSPS activity is defined as the amount of enzyme that catalyses the liberation of 1 mol inorganic phosphate per min reaction time. Determined at 1.0 mM PEP and 1.0 mM S3P.
cDetermined by the Lineweaver-Burk method. The PEP or S3P concentration was fixed at 0.05, 0.075, 0.1, 0.2, 0.5, and 1 mM, respectively, while the other concentration was fixed at 1 mM.
dThe half maximal inhibitory concentration (IC50) is a measure of the effectiveness of glyphosate in inhibiting biochemical function. It was determined by fitting the data to the equation: $V = V_{max} + (V_{max} - V_{min})(1 + (I/C_{50})^b)$, and $V$ was determined at 1 mM PEP and 1 mM S3P with the glyphosate concentration ranging from 0.0001 mM to 100 mM.

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pAroA-Ra5 with about 2.0 kb insert fragment was selected to be sequenced subsequently.

**Gene cloning and mutagenesis**

Primers GR20Z2 (5′-GAGAGAGGATCCATGGAATCCCTCTGAGCATTACAA-3′) and GR20F2 (5′-GAGAGAGGATCCGTGATATGGGATTTGCCTG-3′) were used to amplify the EPSPS gene *AroAR.aquatilis* from pAroA-Ra5. Multiple-site mutagenesis was carried out in order to induce mutations in the hinge region (Thr38Lys and Arg40Val) and the subdomain S5 (Arg292Glu, Ser292Val, Ile225Val, and Gln226Lys) of *AroAR.aquatilis* by PAG-mediated overlap extension PCR [30]. The following oligonucleotide pairs were used: M1Z (5′-GGTTTGGCAGGCTAAACCCTCGTACGATACCTGCTG-3′) and M1F (5′-TCGAGGAGTTCAGTGAGCGGTTTATCCCTGGCTGAAAAGC-3′) for mutations in the hinge region and M2Z (5′-CGGAGCTCTTTGACGACAAATTGCTGGTACTCGTGAGCATGGAACACCAATTTGTCGTTGAAAGAAGCTCTGCAGCAGATTACCCGC-3′) and M2F (5′-TAATGCTGACGCTCTTTGACAATTTGTCGTTGAAAGAAGCTCTGCAGCAGATTACCCGC-3′) for mutations in subdomain 3. The first PCR products were performed with amplified templates combined with the above three purified DNA fragments. Single-site mutagenesis was carried out to obtain the mutants *m3AroAR.aquatilis* with the subdomain 5 of *AroAR.aquatilis* and the mutants *m2AroAR.aquatilis* through direct amplification from pAroA-Ra5 with the primer M2Z and M2F, and M2Z and GR20F. PCR products were resolved using 8% polyacrylamide gels, recovered and purified by electroelution into dialysis bag, and concentrated by ethanol precipitation. The full-length mutant gene *AroAR.aquatilis* with a mutation was obtained by PCR with a pair of primer GR20Z and GR20F and the mixed templates combined with the above three purified DNA fragments. Single-site mutagenesis was carried out to obtain the mutant gene *m5AroAR.aquatilis* or *m3AroAR.aquatilis* through direct amplification from pAroA-Ra5 with the primer MIZ and M1F or MIZ and M2F. In order to further testify that the *f*-strand of the subdomain 5 is important for glyphosate binding, we have generated *m5AroAR.aquatilis* with the subdomain 5 of *AroAR.aquatilis*. Also, the *m5AroAR.aquatilis* and *m5AroAR.aquatilis* were created using a PCR-based staggered extension process.

**Comparison of cell growth in the presence of glyphosate**

Primers EZ1 (5′-CGGGATCCGTTAATGCTGGAATTCTTTCGTTAAAACCGGCCGAC-3′) and EF1 (5′-CGGAGCTCAGGCTCGGAAAAACCGGCCGAC-3′) were used to amplify the EPSPS gene *AroAR.coh* from the chromosomal DNA of *E.coh*. The amplified fragments of the wild type gene *AroAR.coh*, mutant gene *m3AroAR.coh* and *AroAR.coh* were digested with *BamHI* and *SacI* and ligated into the plasmid pYPX251 vector (GenBank accession number AY178046) to construct recombinant plasmids p251_AroA-Ra, p251-m3AroA-Ra and p251-AroA-Ec, respectively. These EPSPS genes were driven by the moderate promoter of *AacC1*.

The *E. coh* aroA-deleted mutant strain ER2799 harboring either p251-AroA-Ra, p251-m3AroA-Ra, and p251-AroA-Ec was grown at 37 °C in liquid M9 minimal medium supplemented with glyphosate at various concentrations ranging from 0 to 150 mM. Cell densities were measured by attenuation at 600 nm [11].

**Construction of plant expression vector and plant transformation**

A DNA fragment encoding the chloroplast transit peptide of tobacco (TSP; GenBank accession number M61904) was PCR amplified from the genomic DNA of tobacco using primers TSPZ (5′-CGGCTTCAGTTGGCAGAGGTAAATCTGGCAGATTG-3′) and TSPF (5′-CAGAGGATCTCTGTGCGGTAGTCCACACTGAT-3′). The PCR fragment was digested with *SalI* and *BamHI*, cloned into the corresponding restriction sites of pGEM3Z (Promega, Madison, WI, USA), and confirmed by DNA sequencing. The *Pst*I fragment of double CaMV35S promoter was inserted upstream of the TSP sequence in the sequence confirmed plasmid. The fused fragment and *aroA* gene were cloned into the Agrobacterium binary vector pYF716 [13] using *Pst*I-BohI and *BamHI*-SacI restriction sites, respectively. The final constructs DSSS:TPsAroANos (Fig. 4A) were introduced into *A. tumefaciens* LBA4404 (Clontech, Palo Alto, CA) by electroporation. The tobacco (*Nicotiana tabacum cv Xanthi*) was transformed by the leaf discs method via the Agrobacterium-mediated transformation procedure described by Wang et al. [32]. Shoots were rooted on a medium containing 30 mg/L hygromycin and then transferred into soil and grown in a greenhouse.

**Transgenic plants selection and assay for herbicide resistance**

The transgenic plants were confirmed by PCR with the specific primers GR20Z2 and GR20F2 for *AroAR.aquatilis*, *m3AroAR.aquatilis* and EZ1 and EF1 for *AroAR.coh*. PCR amplification was performed using the genomic DNA of hygromycin-resistant plants as template. RT-PCR was performed to confirm the transcription of the *AroAR.aquatilis* (*m3AroAR.aquatilis* and *AroAR.coh*), Total RNA was extracted from the seedlings of *T*1 generations using an RNA isolation kit (MBI, Fermentas), and first-strand cDNA was synthesized using 5 μg total RNA with the Reverse Transcription System (Promega, Madison, WI, USA). The fragments of the tobacco Actin gene were amplified from the same cDNAs as a control to normalize the amount of cDNA using primers TactZ (5′-CAATGGACATTTTGCAGTTGGCGCTC-3′) and TactF (5′-CGGAAATCTTCTCGACGCAAATGG-3′). The specific fragment of about 200 bp from *AroAR.aquatilis*, *m3AroAR.aquatilis* and *AroAR.coh* gene was amplified from transgenic plants using the same amount of cDNA. The PCR products were separated on 2% agarose gel and quantified using a Model Gel Doc 1000 (Bio-Rad, USA). The expression pattern of those *aroA* genes was evaluated with a Shine Tech Gel Analyzer (Shanghai Shine Science of Technology Co., Ltd., China). The same results were obtained for three independent experiments.

The leaf discs were cut from individual transgenic tobacco and then moved to Murashige and Skoog (MS) medium containing 0 to 0.8 mM glyphosate and grown for 3 weeks with 16 h of light and 8 h of dark. Seedlings of *T*1 generations were transplanted into soil and grown in five- to six-leaf stage. After spraying with the 5 mM glyphosate for 2 weeks, injury of the transgenic plants was observed visually.

**Overexpression and purification of AroA**

DNA fragments containing the entire coding region of *AroAR.aquatilis* and the mutants *m3AroAR.aquatilis*, *m3AroAR.aquatilis* and *m3AroAR.aquatilis* was obtained by PCR from the positive colony with the following two primers: GR20Z (5′-GAACCATG-GAATCCCTGAGAATTCAAGATCCGTG-3′) and GR20F (5′-GGTTCGAGTGGCAGTGATGTGCTGATCGCGCAGCC-3′). Primer GR20Z contains a mutation where the initiation start codon ATG was replaced with UUG. The fragment was cloned into *NcoI* and *XhoI*, cloned into the corresponding restriction sites of PET-28a (Novagen, Inc.), and confirmed by DNA sequencing. Then the plasmid was transformed to *E. coli* BL21 (DE3; Novagen, Inc.). The expressed protein was purified using a HiTrap HP kit (Amersham Biosciences) according to the manufacturer’s instructions. The AroAR.coh from *E. coli* was overexpressed and purified according to the method used above. Also the protein of *AroAR.aquatilis* was further blotted onto a polyvinylidene difluoride
membrane for N-terminal protein sequencing using an Applied Biosystems Procise sequencer.

Enzyme assay
EPSPS activity was measured by determining the amount of inorganic phosphate produced in the reaction using the malachite green dye assay method [33]. The standard reaction was carried out at 25°C in a final volume of 50 µl containing 30 mM HEPES (pH 7.0), 1 mM S3P, 1 mM PEP, and the purified enzyme. The enzyme was allowed to react for 3 to 5 min before the addition of 800 µl of a 34% sodium citrate solution. After incubation for 30 min at room temperature, the absorbances of the samples were measured at 660 nm. In this case, the same reaction solution without S3P was used as the zero control.

DNA sequence GenBank Accessions
The novel 1284 bp DNA sequence encoding the 5-enolpyruvylshikimate-3-phosphate synthase from *Rahnella aquatilis* was deposited as GenBank Accession number GQ499276.

Supporting Information
Figure S1 Overexpression and purification of *Aro A* from *R. aquatilis*. Lane 1, molecular mass marker; lane 2, protein of *E. coli* ER2799; lane 3, overexpression of *Aro A* from *R. aquatilis* induced with IPTG; lane 4, purified protein with HisTrap HP kit.

Figure S2 Kinetic properties of *Aro A* from *R. aquatilis*. Activities were assayed in Hepes buffer at 28°C in the presence of 1 mM S3P and various concentrations of PEP. A. Steady-state kinetics of *Aro A* from *R. aquatilis* wild type *Aro A* from *R. aquatilis* wild type *Aro A* from *R. aquatilis* wild type Activities were assayed in Hepes buffer at 28°C in the presence of 1 mM PEP and various concentrations of S3P. C. IC50 values of glyphosate inhibition were determined by fitting the data to the equation $v = V_{max} \left(1 - \frac{[I]}{IC50}\right)$, and $v$ was determined at 1 mM PEP and 1 mM S3P, with glyphosate concentrations ranging from 0.001 to 100 mM.

Author Contributions
Conceived and designed the experiments: QHY RHP.Performed the experiments: YST WZ XYF. Analyzed the data: HJH CC XFJ. Contributed reagents/materials/analysis tools: ASX. Wrote the paper: RHP.

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