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Identification and functional characterization of novel plant UDP-glycosyltransferase (*Lb*UGT72B10) for the bioremediation of 3,4-dichloroaniline

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Herbicides are chemicals used to manipulate or control the growth of undesired plants and thereby protect crops. However, they and their degradation products can persist and accumulate in the environment, leading to contamination of soil and water systems and biodiversity loss. Interestingly, through the action of UDP-glycosyltransferases (UGTs), higher plants can glycosylate these xenobiotics, increasing their solubility and alleviating their toxicity. Here, seven plant UGTs belonging to family 72 of the UGT nomenclature were identified to N-glycosylate 3,4-dichloroaniline (3,4-DCA), which is a degradation product of commercially significant herbicides like Diuron, Linuron and Propanil. Although chlorinated chemicals are well-known UGT substrates, only one UGT with activity on 3,4-DCA (AtUGT72B1 from $Arabidopsis\ thaliana$) has been fully biochemically characterized. In this study, biochemical analysis revealed that six of the seven identified UGTs are capable of full conversion of 3,4-DCA to its N-glucoside. The most efficient enzyme was found to be LbUGT72B10 from $Lycium\ barbarum\ (k_{cat}=11.2\ s^{-1},\ K_M=51.2\ \mu\text{M})$. Consequently, transgenic expression of LbUGT72B10 could potentially play a role in the future in the mitigation of 3,4-DCA toxicity, preventing its accumulation in living systems and reducing contamination of waterways and soil.

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Sustainability spotlight

The persistent environmental contaminant 3,4-dichloroaniline (3,4-DCA) results from the degradation of herbicides and industrial pollutants. UDP-glycosyltransferases (UGTs) can facilitate the bioremediation of 3,4-DCA through glycosylation, promoting its biodegradation in the environment and lowering human exposure to the chemical. This study reports the identification and functional characterization of *Lb*UGT72B10, a novel *N*-glycosyltransferase active on 3,4-DCA. Although further investigations are required to fully understand the actual bioremediation mechanism, UGT-mediated glycosylation can help mitigate the health risks associated with the pollutant and improve its biodegradation from water and soil sources, resulting in better water quality and reduced biodiversity loss. Therefore, this study addresses various sustainability challenges aligned with the UN Sustainable Development Goals, including SDG 3, 6, 14, and 15, related to improving human well-being, clean water, preserving life below water, and promoting life on land.

Introduction

Chlorinated xenobiotics are a group of artificially produced chemicals characterized by a cyclic structure and a variable number of chlorine atoms.¹ They are used as crop-protecting agents against weeds and pests or in the manufacturing of pesticides, and they also play a role in the pulp and paper industry.^{2,3} They are classified as Persistent Organic Pollutants (POPs) and, due to their lipophilic nature and persistence, they possess high bioaccumulation potential.⁴⁻⁶ These pollutants can enter the ecosystem *via* a variety of waste streams, but primarily through pesticide application, posing a threat to biodiversity.⁷ Due to their high hydrophobicity, they accumulate

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in the fat tissues of animals and in different structures of plants.^{8,9} Additionally, despite being formulated to target specific categories of weeds, these compounds can be nonselective and cause harm to most plants they come in contact with.10,11 In animals including humans, long-term exposure to these chemicals has been shown to cause neurotoxic effects, liver and kidney damage and possibly cancer.12 Chlorinated anilines are a major class of xenobiotics in the environment.¹³ They are mainly released in the environment as biodegradation products of economically significant herbicides such as Propanil, Diuron and Linuron.14 Particularly, dichloroanilines, such as 3,4-dichloroaniline (3,4-DCA), persist in the environment for a long time as insoluble residues in soil and plant systems, and are detected more frequently compared to their parent compounds. 15-17 3,4-DCA has been detected in surface waters of the European Ebro river basin, with average concentrations ranging from 25 ng L^{-1} to 765 ng L^{-1} . It has also been found in

effluents from dye-manufacturing plants and water field cultures in other parts of the world. 19,20 Notably, high concentrations of 3,4-DCA have been recorded in water samples, reaching up to 567 $\mu g L^{-1}$, and in soil samples from Brazilian rice fields, with levels reaching 119 mg kg⁻¹.²¹ 3,4-DCA is toxic to aquatic organisms and has been shown to induce genotoxic effects, reproductive toxicity and histopathological alteration in mice, being in this way potentially harmful to humans.²² It has also been shown to be more toxic than its parent chemicals.23 Once introduced into the environment, 3,4-DCA is rapidly absorbed by the roots and translocated to the leaves and other parts of the plant.24

Interestingly, plants can regulate the biological activity of pollutants, pesticides, and their metabolites by converting them into less toxic compounds through a four-phase detoxification system: first, xenobiotics are metabolically activated by "phase 1" enzymes, which then facilitates their subsequent bioconjugation with polar natural products in "phase 2" metabolism.25 In crops and weeds, the most observed "phase 2" reaction is glycosylation, which is a biochemical modification catalyzed by a family of enzymes known as UDPglycosyltransferases (UGTs).26 Plant UGTs catalyze the transfer of active sugar groups from nucleotide sugars, usually from uridine diphosphate-glucose (UDP-Glc) to small molecules, such as phytohormones and secondary metabolites, as well as xenobiotics (Fig. 1).27-30 In "phase 3", the glycosides can either be imported into the vacuoles or they can go through additional conjugation with malonyl-CoA, which is a frequent reaction in higher plants that results in the formation of non-phytotoxic secondary conjugates.31,32 A study has shown that the root culture of Arabidopsis thaliana can quickly conjugate 3,4dichloroaniline to 3,4-dichloroaniline-N-glucoside, which is then transported outside the roots.33 The vacuolar storage and the release into the rhizosphere reduce the phytotoxicity of the xenobiotic and prevent yield losses. Therefore, by increasing the water solubility, glycosylation of xenobiotics can help prevent bioaccumulation.32 Interestingly, the model plant Arabidopsis thaliana contains more than 100 UGTs, which are dedicated to small molecule conjugation, and 44 of them catalyze the Oglycosylation of chlorinated chemicals.31 Multiple studies have confirmed that many UGTs can glycosylate them as well as their natural acceptor.34-37 Nonetheless, only a limited number of UGTs, specifically 10, have been discovered to have low Nglycosylation activity towards 3,4-DCA. 13,33,35,36,38 Out of these Nglycosyltransferases (N-GTs), AtUGT72B1 was the only enzyme found to have appreciable N-glycosylation activity, and it was also the only one that has been biochemically characterized with this substrate.31 In the present work, we selected seven candidates from our UGT collection for 3,4-DCA glycosylation, based on sequence identity (>60%) with AtUGT72B1. Of these seven candidates, which belong to the UGT72 family, six showed full conversion of 3,4-DCA to its N-glucoside, and LbUGT72B10 was identified as the most efficient enzyme and biochemically and kinetically characterized. Our in vitro results show that LbUGT72B10 is an efficient biocatalyst for the conversion of the xenobiotic to 3,4-DCA-N-glucoside (Fig. 1). These results highlight the potential value of LbUGT72B10 in

bioremediation strategies aimed at removing the pollutant from the environment by utilizing transgenic technology to manipulate xenobiotic metabolism in plants. However, additional in vivo investigations are required to validate and clarify its effectiveness as a potential bioremediation strategy in order to enhance agricultural sustainability.

2. Materials and methods

Reagents and chemicals

Buffers, standard reagents, and 3,4-dichloroaniline (≥98%) were purchased from Sigma-Aldrich.

2.2 Sequence selection

To identify the closest homologs to the known 3,4-DCAglycosylator AtUGT72B1 (Uniprot Accession: Q9M156, Gen-Bank Accession: AF360262.1) available in our laboratory, the sequence of AtUGT72B1 was used as a query for the NCBI Basic Local Alignment Search Tool (BLASTp) search against our inhouse UGT library sequences.39 Our in-house UGT library is a random selection of UGTs from phylogenetic group E and contains 45 sequences from various plant species, which are readily available from prior studies within our group, as plasmids suitable for the expression of soluble recombinant proteins. The candidate sequences were selected based on their >60% sequence identity with the query sequence. Multiple sequence alignment (MSA) was generated using the Clustal (https://www.ebi.ac.uk/Tools/msa/clustalo/) Omega and visualized with Jalview software (http://www.jalview.org/).

2.3 Cloning, expression and purification of recombinant **UGTs**

The full-length coding sequence of the UGTs was synthesized and cloned into the pET28a(+) vector by Genscript (USA), using the NcoI and XhoI restriction sites. The protein coding sequences included an N-terminal sequence consisting of a 6xHis-tag. E. coli BL21 StarTM (DE3) cells (Thermo Fisher Scientific) were transformed with the plasmids and the precultures were grown overnight at 37 °C in LB medium supplemented with 50 μg mL⁻¹ kanamycin. Main cultures were prepared by adding 5 mL overnight culture to 500 mL of 2xYT culture with 50 μg mL⁻¹ kanamycin and grown at 37 °C and

Fig. 1 Glycosylation reaction of 3,4-DCA to 3,4-DCA-N-glucoside catalyzed by a N-glycosyltransferase (N-GT).

220 rpm until $OD_{600} = 0.5$ –0.7. The cultures were induced with 0.5 mM IPTG and grown for about 22 hours at 20 °C. The cells were harvested by centrifugation at 4500 \times g at 4 °C and resuspended in a buffer containing 50 mM HEPES, 300 mM NaCl, pH 7.5, with 20 mM imidazole supplemented with 0.4 mg DNaseI. The resuspended cells were subjected to cell lysis using an Avestin Emulsiflex C5 homogenizer (ATA Scientific Pty Ltd., Canada), and the soluble cell lysate was recovered by centrifugation at 14 500 \times g at 4 °C for 50 min. The supernatant containing the recombinant proteins was filtered with a 0.22 µM filter and purified by nickel-affinity chromatography (1 mL prepacked HisTrap™ FF columns, GE Healthcare, Sweden) on an ÄKTA pure (GE Healthcare, Sweden). Prior to sample loading, the column was washed and equilibrated with the same buffer (50 mM HEPES, 300 mM NaCl, pH 7.5, with 20 mM imidazole). Following protein binding, the column was washed with a buffer containing 50 mM HEPES, 300 mM NaCl, pH 7.5, with 500 mM imidazole. Elution of the proteins was achieved using a combination of isocratic and gradient elution with an imidazole concentration increasing from 20 to 500 mM. The elution peak fractions were pooled together and concentrated using centrifugal filters (MW cut-off 10-30 kDa), and buffer exchanged to a buffer containing 25 mM HEPES, 150 mM NaCl, pH 7, with 1 mM DTT. The total concentration of partially purified proteins was measured by spectrophotometric measurements at 280 nm in a NanoDrop Instrument (Thermo Fisher Scientific). The sample purity was assessed using SDS-PAGE.

2.4 Glycosyltransferase activity assay toward 3,4-dichloroaniline

Partially purified protein samples were used in 30 μL of total reaction volume to perform *in vitro* glycosylation reactions. The reaction mixture contained 50 μg mL $^{-1}$ of enzymes, 500 μM 3,4-DCA, 2 mM UDP-Glc in a 50 mM phosphate-citrate buffer, pH 7. The control was made with the reaction mixture without the enzyme. The reaction was incubated at 20 °C and quenched at various time intervals (2 min, 5 min and 10 min) by adding 1 volume of ice-cold acetonitrile. Data were analyzed as mean \pm S.D. of two independent experiments.

2.5 Biochemical characterization of LbUGT72B10

For the identification of the pH optimum of LbUGT72B10, its activity was investigated in 100 mM Tris-BisTris buffer, pH 5–10, with 2 μ g mL⁻¹ enzyme, 50 μ M 3,4-DCA and 200 μ M UDP-

Glc. 10 μL of reaction were quenched with 240 μL 0.1% acetic acid at 0, 2, 4, 6, 8 and 10 minutes.

For the identification of the temperature optimum, 2 μg mL $^{-1}$ enzyme activity towards 50 μM 3,4-DCA were assayed in 100 mM Tris–BisTris buffer, pH 7.75, with 200 μM UDP-Glc. The reactions were carried out at 30–54 °C in a thermocycler and were stopped by the addition of 240 μL acetic acid (0.1%) to 10 μL of the reaction mix after 5, 15, 60 and 180 min. In both cases, data were analyzed as mean \pm S.D. of two independent experiments.

For the identification of the melting temperature, differential scanning fluorimetry (DSF) was performed using the Protein Thermal Shift Dye Kit (Thermo Fisher Scientific) and a qPCR QuantStudio5 machine. Dye solution (1000×) was diluted to 2× in 100 mM citrate-phosphate buffer, pH 7. 10 μL of diluted dye solution was mixed with 10 μL of 0.8 mg mL $^{-1}$ LbUGT72B10 in 2× of the same buffer and pipetted into a qPCR 96-wells plate. The plate was centrifuged for 30 s at 1000 rpm and transferred to the qPCR machine. After 2 min incubation at 25 °C, the temperature increased gradually to 99 °C, with a final incubation of 2 min at 99 °C. Measurements were carried out in triplicate. Raw data were analyzed as mean \pm S.D. of three independent experiments with Protein Thermal ShiftTM Software v1.x.

2.6 Kinetic characterization of LbUGT72B10

For the kinetic characterization, based on the optimized reaction conditions, the activity of 10 ng LbUGT72B10 was assayed towards 5–500 μ M 3,4-DCA in 100 mM Tris–BisTris buffer, pH 7.75 at 30 °C. The measurements were carried out under saturating UDP-Glc conditions (1 mM). The reaction was initiated by adding the acceptor substrate and quenched by adding 240 μ L acetic acid (0.1%) to 10 μ L of the reaction mix after 3 min. The μ M of product formed was obtained from the peak area and the corresponding substrate concentration. Data were analyzed as mean \pm S.D. of three independent experiments. $K_{\rm M}$ and $k_{\rm cat}$ values were determined by fitting the initial velocity data using the Michaelis–Menten model and the drc package in Rstudio (Rstudio version 2022.12.0 + 353, RStudio, Inc.).

2.7 HPLC analysis

In all enzymatic assays, the reaction mixture was analyzed by reverse-phase HPLC using an Ultimate 3000 Series apparatus (Thermo Fisher Scientific) and a kinetex 2.6 μ m C18 100 Å 100 \times 4.6 mm analytical column (Phenomenex). Milli-Q water containing 0.1% formic acid and acetonitrile were used as mobile phases A and B, respectively. A combination of isocratic,

Table 1 UGTs selected for this study

Enzyme	Organism	Common name	GenBank accession	UniProt accession
ZmUGT72G3	Zea mays	Maize	EU955500	B6SRY5
SlUGT72B68	Solanum lycopersicum	Tomato	ADI33725	D7S016
LbUGT72B10	Lycium barbarum	Goji	BAG80556	B6EWZ3
FeUGT72B19	Fagopyrum esculetum	Buckwheat	AB909386	A0A0A1H7P3
NtUGT72B82	Nicotiana tabacum	Tobacco	KJ438810	A0A024AX02
PtUGT1	Persicaria tinctoria	Japanese indigo	MF688772	A0A2R2JFJ4
AtUGT72E3	Arabidopsis thaliana	Thale cress	BT030376	O81498

immediate ramp, and gradients at a flow rate of 1 mL min⁻¹ were applied for the separation of 3,4-DCA and 3,4-DCA-Nglucoside: 0-0.5 min, 2% B; 0.5-1.5 min, 35% B; 1.5-3 min, 3580% B; 3-4.2 min, 98% B; 4.2-5 min, 2% B. The analytes were detected at 260 nm. The HPLC data was monitored and quantified via the Chromeleon software (Thermo Fisher Scientific).

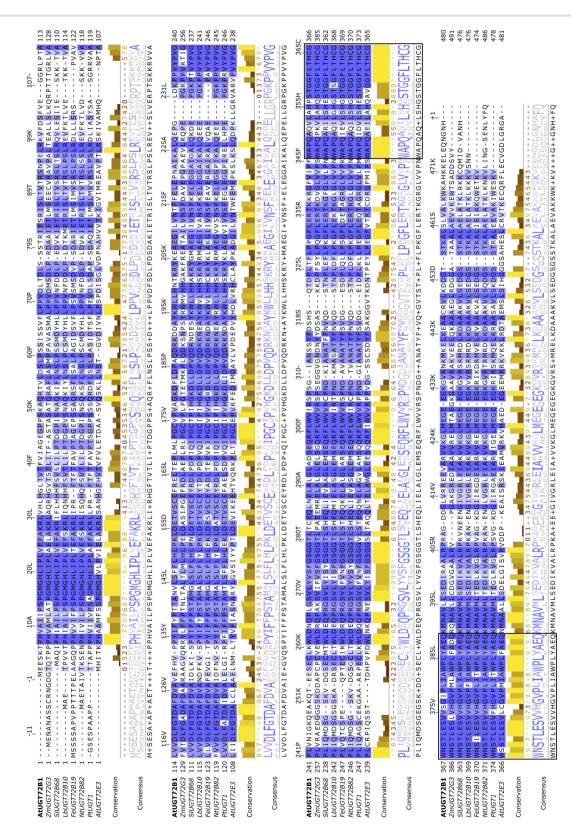


Fig. 2 Multiple sequence alignment of the UGTs selected for this study and AtUGT72B1 (in bold). The PSPG motif characteristic of plant UGTs is boxed.

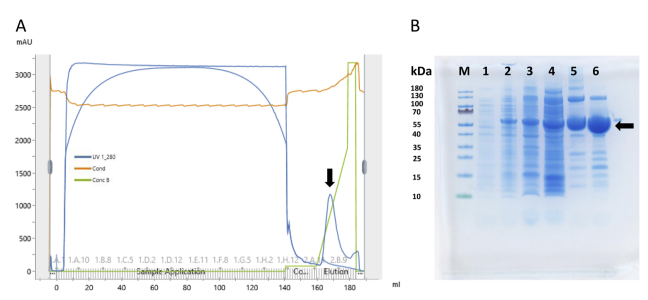


Fig. 3 Purification profile and SDS-PAGE analysis of LbUGT72B10. (A) Nickel-affinity chromatogram of purified LbUGT72B10 using ÄKTA pure (elution peak indicated by the black arrow). (B) SDS-PAGE analysis of the purification steps of LbUGT72B10. (M) PageRuler prestained protein ladder (Thermo Fisher Scientific); (1) E. coli BL21 StarTM (DE3) cells (Thermo Fisher Scientific) before induction with IPTG; (2) E. coli BL21 StarTM (DE3) cells after induction with 0.5 mM IPTG; (3) soluble fraction of clarified E. coli BL21 StarTM (DE3) cell lysate; (4) insoluble fraction of clarified E. coli BL21 StarTM (DE3) cell lysate; (5) purified fraction of LbUGT72B10 following nickel-affinity chromatography purification; (6) purified LbUGT72B10 (indicated by the black arrow) following concentration and buffer exchange, with a predicted MW of \approx 55 kDa.

Results

3.1 Candidate gene selection

To identify additional bioremediation candidates potentially capable of N-glycosylating 3,4-DCA, we selected seven UGT sequences (Table 1) from our "local" enzyme collection with >60% sequence identity to AtUGT72B1 (Uniprot Accession: Q9M156, GenBank Accession: AF360262.1) using NCBI BLASTp search.39 The multiple sequence alignment (MSA) of the selected UGTs and AtUGT72B1 is shown in Fig. 2. All the selected UGTs belong to the UGT72 family, which has been shown to be active with flavonoids, monolignols, chlorophenols, chloroanilines and their precursors or derivatives as substrates.40 The selected UGTs contain the conserved PSPG motif in their C-terminal domain, which is responsible for recognizing the nucleotide-activated sugar (UDP-Glc).41,42 Furthermore, the selected UGTs contain the catalytic histidine believed to play a crucial role in N-glycosylation activity by directing and orienting the nucleophilic attack, without necessarily deprotonating the aniline acceptor. 13,36

3.2 Glycosylation of 3,4-DCA by the recombinantly expressed UGT72 proteins

The selected UGTs were expressed in *E. coli* and purified with nickel-affinity chromatography. The purification results, specifically for *L*bUGT72B10, are presented in Fig. 3. The partially purified proteins were tested for *in vitro* activity toward 3,4-DCA. The enzymatic reaction was followed by reverse-phase HPLC and, since an analytical standard for 3,4-DCA-*N*-glucoside was unavailable, the product of *At*UGT72B1 with 3,4-DCA as substrate was used as reference. With the exception of *At*UGT72E3, all of the enzymes showed *N*-glycosylation activity toward 3,4-DCA.

Remarkably, *Lb*UGT72B10 demonstrated superior performance, converting almost 50% of 3,4-DCA into 3,4-DCA-*N*-glucoside within 2 minutes, and achieving nearly 100% conversion yield after 10 minutes (Fig. 4). For this reason, this enzyme was selected for further biochemical and kinetic characterization.

3.3 Biochemical and kinetic characterization of *Lb*UGT72B10

The pH and temperature profiles of *Lb*UGT72B10 were determined with 3,4-DCA as acceptor and UDP-Glc as donor

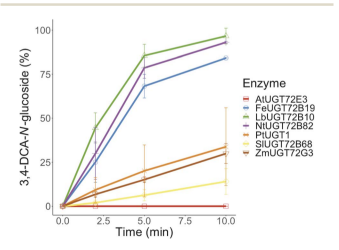


Fig. 4 Glycosylation of 3,4-DCA over time (0–10 min) catalyzed by the expressed recombinant UGTs. Error bars represent standard deviations from the average of duplicate measurements. 50 μ g mL⁻¹ of each purified enzyme was assayed with 0.5 mM 3,4-DCA and 2 mM UDP-Glc in a 50 mM citrate-phosphate buffer, pH 7. Reactions were quenched at different time points and analyzed by reverse-phase HPLC.

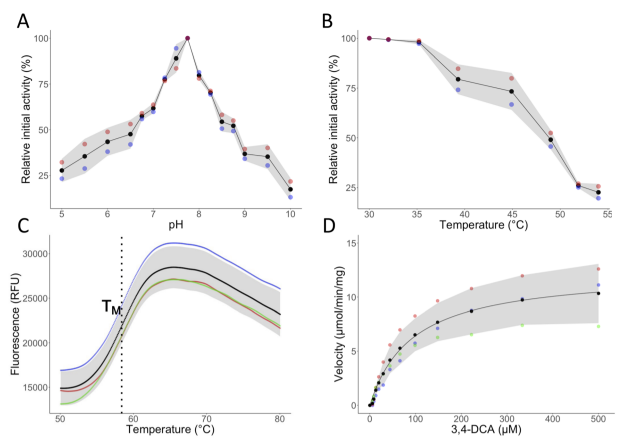


Fig. 5 Biochemical and kinetic characterization of LbUGT72B10. (A) The effect of different pH values on the initial rates of 2 μ g mL⁻¹ LbUGT72B10 in the presence of 50 μM 3,4-DCA at 20 °C in a 100 mM Tris-BisTris buffer (pH 5-10). (B) The effect of different temperatures on the initial rates of 2 μ g mL⁻¹ LbUGT72B10 in the presence of 200 μ M 3,4-DCA in a 100 mM Tris-BisTris buffer, pH 7.75. (C) Thermal stability measured by DSF of 0.8 mg mL⁻¹ LbUGT72B10 in 50 mM citrate-phosphate buffer, pH 7.75. (D) Michaelis-Menten curve for the synthesis of 3,4-DCA-N-glucoside after 3 minutes in the presence of 0.2 ng μ L⁻¹ enzyme at 30 °C, 100 mM Tris-BisTris buffer, pH 7.75. In all the plots, the shaded region represents the standard deviation derived from multiple measurements. Black dots indicate averages from multiple experiments (represented by blue, red and green dots)

substrates (Fig. 5A and B). LbUGT72B10 was found to have a broad pH tolerance, with activity detected in the pH range of 5.0-9.5 and an optimum pH of 7.75. This result agrees with the published values reported for other UGTs, which are in the range 7.5-9.0.43-48 The temperature optimum was found to be 30 °C, with activity rapidly decreasing beyond 35 °C and no detectable activity at 55 °C. This result conforms to the published values for other UGTs, with their optimal temperature range between 30-40 °C.47-50

We further investigated the thermostability of LbUGT72B10 (Fig. 5C). The apparent melting temperature $(T_{\rm M})$ of UGT72B10 was estimated to be approximately 58.4 \pm 0.4 $^{\circ}\text{C}$ using Differential Scanning Fluorimetry (DSF). This is remarkably high for a UGT, which normally melts at approximately 45 °C.37 Although no activity was found beyond 55 °C, this relatively high melting temperature might indicate that LbUGT72B10 possesses superior temporal and chemical stability compared to other UGTs, since these properties often go hand in hand. Our previous study has shown that the reaction conditions to which GT1 enzymes are exposed can impact their thermal stability.37 However, it should be noted that changes in one kind of stability might have an impact on other aspects of stability as well. For example, alterations in pH

or temperature can impact conformational and chemical stability, while chemical modifications or denaturation can affect thermal and conformational stability.51-56

After determining the optimal pH and temperature, we proceeded to performing kinetic characterization (Fig. 5D and Table 2). LbUGT72B10 exhibited a comparable catalytic efficiency to that of the previously reported AtUGT72B1.13 However, its k_{cat} is 3-fold higher than that of AtUGT72B1 and 10-fold higher than that of PtUGT1.13,36 On the other hand, the KM of LbUGT72B10 is around three orders of magnitude higher than that of AtUGT72B1.¹³ Nevertheless, this affinity constant is still considered relatively low when compared to other members of the UGT72 family acting on various acceptors. 47,57,58

Table 2 Apparent catalytic values of LbUGT72B10 and previously reported 3,4-DCA N-GTs

Enzyme	$K_{\mathbf{M}}\left(\mu\mathbf{M}\right)$	$k_{\rm cat} ({\rm s}^{-1})$	$k_{\text{cat}}/K_{\text{M}} \left(\text{M}^{-1} \text{ s}^{-1} \right)$	Reference
LbUGT72B10 AtUGT72B1	51.2 ± 24.2 16.5 ± 4.0		2.33×10^{5} 2.54×10^{5}	This study
PtUGT1	—	1.2 ± 0.3 1.2 ± 0.1		36

4. Discussion

Previous studies have suggested that transgenic plants overexpressing specific glycosyltransferases could serve as effective tools for bioremediation. 33,38,59 However, our understanding of UGTs involved in the N-glycosylation of 3,4-DCA remains limited, both in vivo and in vitro, and their characterization is lacking. Therefore, this study aimed at expanding our knowledge of UGT-mediated glycosylation of the persistent pollutant 3,4-DCA, since this biological reaction could serve as tool in bioremediation efforts, with the view to expand arable land area and thereby enhance global food security. 17 Based on sequence similarity (>60%) to AtUGT72B1, a known glycosylator of 3,4-DCA, we selected seven novel UGT candidates to test for enzymatic activity toward 3,4-DCA. All chosen UGTs had the conserved PSPG motif and the catalytic histidine necessary for N-glycosylation specificity. 13,36 All enzymes except AtUGT72E3 showed N-glycosylation activity within 10 minutes, with LbUGT72B10 reaching approximately 100% yield after 10 minutes. LbUGT72B10 was further biochemically and kinetically characterized. The enzyme was active in pH from 5.0 to 9.5, with an optimum pH at 7.75 and a temperature optimum of 30 $^{\circ}$ C. LbUGT72B10 had a remarkably high melting temperature (58.4 °C), suggesting that the enzyme also possesses chemical stability, as these properties are often interconnected.^{37,54} These characteristics, together with excellent kinetic properties (full conversion in 10 minutes, $K_{\rm M}=51.2~\mu{\rm M}$ and $k_{\rm cat}=11.2~{\rm s}^{-1}$ align well with the requirements for enzymes employed in various bioremediation strategies. 60,61

The potential bioremediation approach using transgenic expression of LbUGT72B10 depends on the reproducibility of these in vitro results under in planta conditions, taking into account the influence of the cellular environment on enzyme activity and stability. Also, it is worth noting that currently no genetically modified crops with UGT genes are in use, despite the existence of patented plant UGT sequences for crop protection and bioremediation applications.30 However, even if the in vitro results do not precisely reflect the in vivo context, transgenic plants expressing LbUGT72B10 may still exhibit enhanced resilience toward 3,4-DCA, potentially showing enhanced cellular compartmentalization due to glycosylation, and/or via the higher water solubility of the glycoside promoting efficient 3,4-DCA excretion into the soil, where it can be mineralized by soil microbes.32 3,4-DCA is primarily metabolized through catabolic pathways into naturally occurring intermediates, such as catechol.62 However, only specific microorganisms have been shown to possess 3,4-DCA resistance and to be able to metabolize high concentrations of this xenobiotic, so glycosylation could also mitigate its toxicity toward naturally sensitive microbial populations. 63 Pichia pastoris, a yeast commonly used in biochemical research which is naturally sensitive to 3,4-DCA, has been shown to survive in the presence of this compound when a UGT, VvUGT72B1, was heterologously expressed in the yeast cells.64 A similar phenomenon may occur in soil microbial populations, if the glucoside rather than the aglycon is excreted in the rhizosphere.

Nevertheless, further investigations are required to elucidate the actual bioremediation mechanism in transgenic plants overexpressing *Lb*UGT72B10 and to determine the fate of 3,4-DCA-*N*-glucoside and its metabolism by soil microorganisms.

5. Conclusions

In conclusion, we have studied seven novel UDP-glycosyltransferases for the glycosylation of the recalcitrant pollutant 3,4-DCA. Among them, *Lb*UGT72B10 from *Lycium barbarum* exhibited the highest *N*-glycosylation activity, as well as superior stability and excellent kinetic properties. Although further investigations are needed, the identification of *Lb*UGT72B10 as an efficient enzyme for the glycosylation of 3,4-DCA with potentially superior stability makes it interesting for potential future use in bioremediation efforts of contaminated environments, with the view to expand arable land area and enhance food security and agricultural sustainability.

Conflicts of interest

There are no conflicts to declare.

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