Recruitment of archaeal DTD is a key event toward the emergence of land plants

Mohd Mazeed1,*, Raghvendra Singh1,*, Pradeep Kumar1,2, Ankit Roy1, Bakthisaran Raman1, Shobha P. Kruparani1, Rajan Sankaranarayanan1,2†

Streptophyte algae emerged as a land plant with adaptations that eventually led to terrestrialization. Land plants encounter a range of biotic and abiotic stresses that elicit anaerobic stress responses. Here, we show that acetaldehyde, a toxic metabolite of anaerobic stress, targets and generates ethyl adducts on aminoacyl-tRNA, a central component of the translation machinery. However, elongation factor thermo unstable (EF-Tu) safeguards L-aminoacyl-tRNA, but not D-aminoacyl-tRNA, from being modified by acetaldehyde. We identified a unique activity of archaeal-derived chiral proofreading module, D-aminoacyl-tRNA deacetylase 2 (DTD2), that removes N-ethyl adducts formed on D-aminoacyl-tRNAs (NEDATs). Thus, the study provides the molecular basis of ethanol and acetaldehyde hypersensitivity in DTD2 knockout plants. We uncovered an important gene transfer event from methanogenic archaea to the ancestor of land plants. While missing in other algal lineages, DTD2 is conserved from streptophyte algae to land plants, suggesting its role toward the emergence and evolution of land plants.

INTRODUCTION

The root is a critical architectural innovation in land plants, which is essential for anchorage, nutrient, and water uptake (1, 2). Plants being sedentary in their lifestyle, roots have to survive in the oxygen-deprived environment of soil and face multiple stresses, including major anaerobic stress. Thus, roots are the primary sites of acetaldehyde production, an intermediate of ethanol fermentation (3–7). Environmental assaults such as floods, waterlogging, and parasite infection worsen the situation by creating prolonged anoxic conditions, which eventually produce excess acetaldehyde (3, 4, 8). Acetaldehyde is also an ever-present metabolite in plants and involved in multistress response. Moreover, important processes such as seed germination and pollen development are highly dependent on ethanol fermentation. Anomalies in this pathway lead to failure in seed vigor and male fertility (8–12).

Along with anaerobic stress, roots experience tremendous stress from d-amino acids that are found in abundance in the soil. However, the presence of d-amino acids in plants is inevitable because they play vital roles in plant physiology, such as in development of pollen tubes, apart from acting as a primary nitrogen source (13). However, d-amino acids also cause cellular toxicity by generating d-aminoacyl-tRNAs (d-aa-tRNAs), which sequester a free tRNA pool and impede translation. Removal of mischarged d-amino acids from d-aa-tRNAs is termed chiral proofreading, which enables the perpetuation of homochirality in the cellular proteome (14, 15). d-amino acid editing modules are universally conserved; d-aa-tRNA deacetylase 1 (DTD1) is found in both Bacteria and Eukarya, and DTD2 is present in Archaea and Plantae. Earlier, the presence of DTD2 was noted in all the sequenced genomes of the plant kingdom available at that time. Hence, it was assumed that DTD2 is conserved throughout the plant kingdom (Archaeprotista) (16–18). Plants therefore are unique in having both of the evolutionarily distinct chiral proofreading modules DTD1 and DTD2. Unexpectedly, DTD2 knockout plants, using Arabidopsis as a model system, have been shown to display pronounced sensitivity to both ethanol and acetaldehyde, which are metabolites generated by sustained ethanol fermentation (19, 20).

Acetaldehyde is one of the highly reactive metabolites present inside the cell. It is known to be an unavoidable genotoxic intermediate of anaerobic fermentation. Acetaldehyde forms adducts with DNA and proteins by making an electrophilic attack on free amino groups (21–25). The toxic effects of acetaldehyde are quenched by multiple tiers of detoxifying systems. The first line of defense is provided by acetaldehyde dehydrogenase (ALDH), which converts acetaldehyde to nonreactive acetate. The second level of protection is offered by DNA repair mechanisms such as Fanconi anemia group D2 protein (FANCD2), which maintains chromosomal integrity and stability by repairing acetaldehyde-driven DNA damage. In alcoholic mice and humans, failure of acetaldehyde surveillance mechanisms leads to cancers such as Fanconi anemia, esophageal cancer, and hepatocellular carcinoma (26, 27). Catastrophic effects of acetaldehyde have been reported in various organisms, including plants (28–30). The most intriguing finding among plants is the acetaldehyde hypersensitivity of DTD2 knockouts (17, 19, 20). It has been a longstanding question how a d-amino acid detoxifying system DTD2 confers protection to plants from toxic intermediates of ethanol fermentation, especially from acetaldehyde during anaerobic stress.

Our study aimed at demystifying the link between DTD2 and its role in protecting plants from acetaldehyde-mediated toxicity. The work provides the first empirical evidence that acetaldehyde targets aa-tRNAs, which are highly sensitive compared to nucleic acids and proteins. Acetaldehyde creates ethyl modification of amino acids of aa-tRNA, which makes aa-tRNA more alkali-stable than peptidyl-tRNA. We further identified a unique role of DTD2 as a proofreader of d-aa-tRNA adducts since neither DTD1 nor peptidyl-tRNA hydrolase (PTH) can recycle N-ethyl-aa-tRNAs (NEATs). The unique activity of DTD2 necessitated its recruitment from Archaea to land plants. This is the first evidence of
archaeal gene transfer from methanogens to streptophyte algae. Moreover, the distribution of DTD2 is strongly correlated with acetaldehyde biosynthesis. The presence of DTD2 in land plants and their ancestors, starting from early-diverging subaerial streptophyte algae, suggests that acquiring DTD2 from Archaea is one of the key events in the emergence of land plants.

RESULTS

Acetaldehyde creates ethyl modification on aa-tRNA

DTD2 is known to proofread D-aa-tRNAs as a part of translational quality control (16, 17). Therefore, we set out to investigate whether DTD2’s ability to protect plants from acetaldehyde has the involvement of aa-tRNAs. Hence, we treated aa-tRNAs with acetaldehyde, followed by a reducing agent (Fig. 1A). Unexpectedly, this resulted in a differential migration of the spot corresponding to aa-tRNA on thin-layer chromatogram (TLC) when compared with untreated control (Fig. 1B). Additional moiety attached to the α-amino group of aa-tRNA, such as a peptidyl or an acetyl group, is known to increase the stability of the ester linkage between the amino acid and the tRNA and is best assessed by alkali treatment (31). Earlier studies have shown that peptidyl-tRNAs and N-acetyl-α-aa-tRNAs (substrate mimics of peptidyl-tRNAs) are more stable compared to aa-tRNAs and are hydrolyzed by PTH (32). Upon performing the alkali treatment experiment, we found that the ester linkage of acetaldehyde-treated aa-tRNA (modified D-Phe-tRNA<sub>Phe</sub>; half-life: ~12 min) showed enhanced stability compared to aa-tRNA (D-Phe-tRNA<sub>Phe</sub>; half-life: ~1 min) and N-acetyl-aa-tRNA (N-acetyl-D-Phe-tRNA<sub>Phe</sub>; half-life: ~2.5 min) (Fig. 1C). Ultrastability of N-blocked aa-tRNAs makes them more lethal to the cell because they permanently arrest the tRNA pool and prevent it from participating in translation, which is evident in the case of a temperature-sensitive strain of Escherichia coli for PTH (32, 33). Acetaldehyde-treated aa-tRNAs are therefore expected to be more toxic to the cell than N-acetyl-α-aa-tRNAs and peptidyl-tRNAs because of their high alkali-stable nature. These observations based on TLC and alkali treatment indicated that acetaldehyde could modify aa-tRNA.

To characterize the chemical nature of the modification, electrospray ionization mass spectrometry (ESI-MS) of both acetaldehyde-treated and untreated nonhydrolyzable analogs of aa-tRNAs was performed. These analogs mimic both amino acid and tRNA parts of aa-tRNA, where the amino acid is bound to adenosine (terminal adenosine-76 of tRNA) with nonhydrolyzable amide bond (14). MS data clearly showed that acetaldehyde covalently modifies aa-tRNA with “ethyl” moiety (Fig. 1D and fig. S1, A and B).

Acetaldehyde modifies amino acid of α-aa-tRNA and is independent of chirality and side chain

Incubation of acetaldehyde with α-aa-tRNAs (Tyr-tRNA<sub>Tyr</sub>, Phe-tRNA<sub>Phe</sub>, Ala-tRNA<sub>Ala</sub>, Ser-tRNA<sub>Thr</sub>, and Thr-tRNA<sub>Thr</sub>) or with their analogs (Tyr2AA, Ala3AA, Val3AA, and Thr2AA) bearing amino acids having either chirality (α- or d-) and varied side chains revealed that acetaldehyde modifies aa-tRNA irrespective of its chirality and side-chain chemistry (Figs. 1B and 2A and figs. S1, C to J, and S2). Since acetaldehyde is known to target free amino groups, to identify the site of ethyl modification on the aa-tRNA, tandem MS fragmentation (MS<sup>2</sup>) of ethyl-modified analogs was carried out. Fragmentation experiments (MS<sup>2</sup>) confirmed that modification happens only on the amino acid part of aa-tRNA (Fig. 2, B to D, and figs. S1 and S2) and not on tRNA. These experiments established the link between acetaldehyde, an anaerobic stress metabolite, and aa-tRNA, a key constituent of the translation machinery (TM). It also prompted us to hypothesize that DTD2 relieves anaerobic stress associated with TM in plants by removing ethyl-modified amino acids from NEATs.

DTD2 alleviates anaerobic stress in plants by recycling NEDATs

To explore whether DTD2 deacylates NEATs, in vitro deacylation assays were performed by incubating DTD2 of Arabidopsis thaliana (At DTD2) with both N-ethyl-α-aa-tRNAs [NELATs; N-ethyl-α-Phe-tRNA<sub>Phe</sub> (At) and N-ethyl-α-Tyr-tRNA<sup>Tyr</sup> (Tht) (<i>Thermus thermophilus</i> (Tht))] and N-ethyl-α-aa-tRNAs [NEDATs; N-ethyl-α-Phe-tRNA<sub>Phe</sub> (At) and N-ethyl-α-Tyr-tRNA<sup>Tyr</sup> (Tht)]. In line with our hypothesis, At DTD2 readily deacylated NEDATs to N-ethyl-α-amino acids and free tRNAs but not NELATs (Fig. 3, A and B, and fig. S3, A and B). The activity of DTD2 on NEDATs is hereafter referred to as “NEDAT deacylase (ND) activity.” Since plants also have the canonical chiral proofreading module DTD1 alongside DTD2, we carried out deacylation of NEATs and NEDATs with DTD1. Our biochemical data revealed that neither bacterial (<i>E. coli</i>) nor eukaryotic (<i>A. thaliana</i>) DTD1 acts on NELATs or NEDATs (Fig. 3, C and D, and figs. S3, C to J, and S4, A and B). DTD1’s weak activity on NEATs is corroborated by the earlier structural studies on DTD1 in complex with the D-aa-tRNA analog d-Tyr3AA (14, 34). The crystal structure had revealed that the α-amino acid of a D-aa-tRNA fits snugly into the active site of DTD1, thereby sterically excluding the N-ethylated amino group of a D-amino acid (Fig. 3E). PTH, as mentioned before, acts on N-acetyl-α-aa-tRNAs, which are similar to ethylated substrates (Fig. 3F). Moreover, PTH and DTD2 share a similar fold (16); therefore, we expected that PTH could act on NEATs. However, to our surprise, neither bacterial (<i>E. coli</i>) nor archaeal and the eukaryotic version of PTH (<i>Sulfolobus tokodaii</i>) had activity on NEATs (Fig. 3, G to J, and fig. S4, C to L), suggesting their strict chemical selectivity. In light of these findings on the biochemical activities of the editing modules DTD1, PTH, and DTD2, DTD2 stands out with a distinct role as a unique proofreader of D-aa-tRNA adducts.

Elongation factor thermo unstable protects α-aa from acetaldehyde

As mentioned above, the biochemical experiments were performed in vitro, and to show the accumulation of NEATs in vivo, we used a surrogate (<i>E. coli</i>) system. Initially, <i>E. coli</i> cells were grown in the presence of D-amino acids and acetaldehyde. Later, aa-tRNAs were probed using Northern blotting. As expected, the aa-tRNAs isolated from bacteria exposed to both D-amino acids and acetaldehyde were stable upon alkali treatment. At the same time, the control samples were less stable (Fig. 4, A to D). To further validate the identity of the alkali-stable aa-tRNAs, we performed MS experiments. MS experiments confirmed that these aa-tRNAs were modified to NEATs by acetaldehyde (Fig. 4E). Similar to in vitro experiments, these in vivo generated NEATs were deacylated by DTD2 (Fig. 4F). However, acetaldehyde also modified 1-aa-tRNAs as efficiently as D-aa-tRNAs in vitro (Figs. 1B and 2A and figs. S1, S2, and S4D). Therefore, it is important to know how 1-aa-tRNAs are protected from acetaldehyde in vivo. Elongation factor thermo
unstable (EF-Tu) is one of the most abundant proteins of the cell and is known to bind \( l \)-aa-tRNAs with higher affinity compared to \( d \)-aa-tRNAs (35). \( d \)-aa-tRNAs are discriminated by the EF-Tu and other cellular chiral checkpoints (36). To test EF-Tu’s ability to protect \( l \)-aa-tRNAs from freely diffusing acetaldehyde, we performed modification experiments in the presence of EF-Tu. These assays clearly showed that EF-Tu offers protection to \( l \)-aa-tRNAs (\( l \)-Phe-tRNA\(^{\text{Phe}} \) and \( l \)-Tyr-tRNA\(^{\text{Tyr}} \)) but not to \( d \)-aa-tRNAs (\( d \)-Phe-tRNA\(^{\text{Phe}} \) and \( d \)-Tyr-tRNA\(^{\text{Tyr}} \)) from acetaldehyde modification (Fig. 5, A to C).

EF-Tu protection experiments led to the identification of its unique role in specifically safeguarding \( l \)-aa-tRNAs from reactive metabolites of the cell such as acetaldehyde apart from delivering them to the ribosome. From the crystal structure, it was evident that the \( \alpha \)-NH\(_2\) group of aa-tRNA is engaged in interactions with EF-Tu, and there is no extra space to accommodate an ethyl group (Fig. 5D) (37).

The ability of DTD2 to rescue acetaldehyde toxicity in Arabidopsis in combination with its lack of activity on NELATs highlights that NEDATs are the physiological substrates of DTD2. Together, the
above findings suggest that DTD2 alleviates the effect of acetaldehyde by actively clearing NEDATs during anaerobic stress.

**ND activity is rooted in the Archaea**

Apart from plants, DTD2 is also conserved in Archaea (Fig. 6A and fig. S5, A and B) (16, 17), and therefore, we further checked whether archaeal DTD2s have the ND activity. We performed NEAT deacylation experiments with distantly related archaeal DTD2s from the organisms *Pyrococcus horikoshii* (*Pho* DTD2), *Methanocaldococcus jannaschii* (*Mj* DTD2), and *Archaeoglobus fulgidus* (*Af* DTD2). All three archaeal DTD2s acted on NEDATs with similar efficiency as that of plant DTD2s (Fig. 6, B to D, and fig. S5C). Similar to plant DTD2s, archaeal DTD2s also did not act on NELATs, suggesting that not only the ND activity but also chiral selectivity toward adducts is conserved in DTD2s of Archaea (Fig. 6E and fig. S5D). The conservation of DTD2 and its ND activity across Archaea highlights that DTD2’s ability to clear NEDATs is “rooted” in the archaeal branch of life. Therefore, it is clear that plants specifically acquired DTD2 from Archaea during evolution. This is further supported by
Fig. 3. NEDATs are readily acted upon by A. thaliana DTD2 but not by DTD1 or PTH. (A) Deacylation of N-ethyl-α-Phe-tRNA\(^{\text{Phe}}\) (At) with At DTD2. (B) Deacylation of N-ethyl-λ-Phe-tRNA\(^{\text{Phe}}\) (At) with At DTD2. (C) Deacylation of N-ethyl-α-Phe-tRNA\(^{\text{Phe}}\) with At DTD1 and Ec DTD1. (D) Deacylation of N-ethyl-λ-Phe-tRNA\(^{\text{Phe}}\) with At DTD1 and Ec DTD1. (E) Crystal structure of DTD1 in complex with α-Tyr3AA. Left of the zoomed-in section of the active site shows the binding mode of the snugly fit substrate; right shows a steric clash between the modeled ethyl modification on the α-NH\(_2\) group of α-amino acid and active site residues. PDB, Protein Data Bank. (F) Schematics showing subtle chemical differences among peptidyl-tRNA, N-acetyl-aa-tRNA, and NEAT (N-acetyl-aa-tRNAs differ with NEATs only by carbonyl oxygen present on the carbon that is attached to the α-NH\(_2\) group of amino acid of aa-tRNA). (G) Deacylation of N-ethyl-α-Phe-tRNA\(^{\text{Phe}}\) (Ec) with Ec PTH. (H) Deacylation of N-ethyl-λ-Phe-tRNA\(^{\text{Phe}}\) (Ec) with Ec PTH. (I) Deacylation of N-ethyl-α-Phe-tRNA\(^{\text{Phe}}\) (Pho) with St PTH2. (J) Deacylation of N-ethyl-λ-Phe-tRNA\(^{\text{Phe}}\) (Pho) with St PTH2. In all our biochemical assays, the concentration of substrate used was 0.2 μM. At, A. thaliana; Ec, E. coli; St, S. tokadii.
Fig. 4. Acetaldehyde generates ethyl modification on d-aa-tRNA in vivo. (A) CuSO₄ + tris (8.0) treatment of in vitro generated d-Phe-tRNA⁰_d, d-Tyr-tRNA⁰_Tyr, modified (N-ethyl)-d-Phe-tRNA⁰_Phe, and modified (N-ethyl)-d-Tyr-tRNA⁰_Tyr. (B) Schematic showing the method used to generate NEDATs in E. coli. (C) Northern blot analysis showing no accumulation of Tyr-tRNA⁰_Tyr adducts in E. coli cells grown in the presence of only d-tyrosine. (D) Northern blot analysis showing the accumulation of Tyr-tRNA⁰_Tyr adducts in E. coli cells grown in the presence of d-tyrosine and acetaldehyde. (E) Alkali hydrolysis (incubation in 15% of NH₄OH at 70°C for 18 hours) of aa-tRNA isolated from Δtdt E. coli, Δtdt dE. coli + d-Tyr (grown in the presence of 3 mM d-tyrosine), and Δtdt dE. coli + 0.01% acetaldehyde (grown in the presence of 0.01% acetaldehyde) yielded tyrosine peaks. N-ethyl-tyrosine peaks were seen only in Δtdt E. coli + 3 mM d-Tyr + 0.005% acetaldehyde (grown in the presence of 3 mM d-tyrosine and 0.005% acetaldehyde), and Δtdt dE. coli + 3 mM d-Tyr + 0.01% acetaldehyde (grown in the presence of 3 mM d-tyrosine and 0.01% acetaldehyde). Fragmentation of the respective peaks also confirms the same. (F) Northern blot analysis showing deacylation of in vivo generated NEDATs by both archaeal DTD2 (Pho DTD2, 50 nM) and plant DTD2 (At DTD2, 50 nM).
Absence in Bacteria and Opisthokonta (Archaeplastida) delineated that DTD, i.e., DTD2. Extensive bioinformatic analysis of DTD2 distribution is exclusive to land plants (Bryophyta, Pteridophyta, Gymnosperms, and Angiosperms) and streptophyte algae (Chlorokybophyceae, Klebsormidiophyceae, Charophyceae, and Zygmenatophyceae). At the same time, DTD2 is entirely absent in different clades of algae, which include glaucophytes, red algae, and green algae. DTD2 is present in the recently sequenced genomes of the streptophyte algae Chlorokybus atmophyicus (42), Klebsormidium nitens (43), Chara braunii (44), and Mesotaenium endlicherianum (45) (Fig. 7A), of which C. atmophyicus is an early-diverging “subaerial/terrestrial” algae (42). Unexpectedly, the early-diverging “aquatic” streptophyte algae Mesostigma viride does not encode DTD2. Structure-based alignment of DTD2 sequences revealed that DTD2 of streptophyte algae have conserved residues, which are essential for catalysis (Fig. 7B). To further validate our sequence analysis, we performed biochemical assays using DTD2 of streptophyte algae [K. nitens DTD2 (Kn DTD2)]. We found that Kn DTD2 not only showed ND activity (Fig. 7C and fig. S5G) but also conserved chiral selectivity (Fig. 7D and fig. S5H) and ethyl specificity on adducts (Fig. 7E and F, and fig. S5, I and J). These observations suggest that DTD2 may have provided an adaptive advantage to streptophyte algae, allowing them to venture on to the land.

**Acquisition of DTD2 is one of the first steps on the long road of land plant evolution**

Recent evidences have shown that streptophyte algae acquired many land plant–specific characteristics such as phytohormone biosynthesis, phytohormone signaling, xylolucan biosynthesis, phytochromes, and cyclic electron flow (42–46). These genes are conserved across land plants and are mostly absent in other clades of algae (42, 43). Similarly, the presence of DTD2 from streptophyte algae to land plants suggests that its acquisition is an important land plant–specific adaptation. In addition, the remarkable conservation...
Fig. 6. DTD2’s ND activity and chiral selectivity are conserved throughout Archaea. (A) Phylogenetic classification of archaeal and plant DTD2s (generated with scoring matrix LG+RS (according to the BIC score), of 546 protein models, LG+RS was chosen as the best amino acid substitution model for the given multiple sequence alignment, which was used to construct a phylogenetic tree). (B) Deacylation of N-ethyl-d-Phe-tRNA\textsuperscript{Phe} (Pho) with Pho DTD2. (C) Deacylation of N-ethyl-o-Phe-tRNA\textsuperscript{Phe} (Pho) with Mj DTD2. (D) Deacylation of N-ethyl-o-Phe-tRNA\textsuperscript{Phe} (Pho) with Af DTD2. (E) Deacylation of N-ethyl-l-Phe-tRNA\textsuperscript{Phe} (Pho) with Pho DTD2, Mj DTD2, and Af DTD2 showing no activity even at 500 nM enzyme concentration. (F) Deacylation of N-acetyl-o-Phe-tRNA\textsuperscript{Phe} with both archaeal (Pho) and plant (At) DTD2s showing no activity even at 500 nM enzyme concentration. (G) Deacylation of N-acetyl-l-Phe-tRNA\textsuperscript{Phe} with both archaeal (Pho) and plant (At) DTD2s showing no activity even at 500 nM enzyme concentration. (H) Co-occurrence of DTD2 with acetaldehyde biosynthesis in Archaea and plants.
Fig. 7. DTD2 is conserved in land plants and streptophyte algae in the plant kingdom. (A) DTD2 distribution is seen only in Archaea, land plants, and streptophyte algae (genome sequences of Coleochaetophyceae members are not available, and hence, DTD2’s status is not depicted in the figure). (B) Structure-based sequence alignment of DTD2 shows conservation of active site residues (16) throughout Archaea, streptophyte algae, and land plants. Deacylation of N-ethyl-d-Phe-tRNA_Phe (At) (C), N-ethyl-l-Phe-tRNA_Phe (At) (D), N-acetyl-d-Phe-tRNA_Phe (At) (E), and N-acetyl-l-Phe-tRNA_Phe (At) (F) with K. nitens DTD2. (G) DTD2 is the only gene of exclusive archaeal origin that was recruited for the adaptation of land plants. (H) Sequence identity values between representative plant DTD2s with all the DTD2 sequences (includes both Archaea and plants) [M. conradii (M), C. atmophyticus (C), and K. nitens (K)] are highlighted in the figure [the horizontal axis represents archaeal DTD2 sequences (1 to 149) and plant DTD2 sequences (150 to 234), and the vertical axis represents sequence identity values]. (I) Boxplot displays DTD2 of M. conradii that shares the maximum identity (~33%) with DTD2s of all the plants; the horizontal axis represents sequence identities with all plant DTD2s, and the vertical axis shows the representative archaeal sequences compared with all plants.
of DTD2 starting from the early-diverging subaerial/terrestrial streptophyte algae *C. atmophyticus* indicates that recruitment of DTD2 is one of the key first steps on the long road of land plant evolution.

Moreover, we searched for the origins of the genes mentioned above that are responsible for the adaptations of land plants. Among such genes, DTD2 is the only gene that has an exclusive archaeal origin. In contrast, all the other genes have either only bacterial origin or mixed origins (both bacterial and archaeal) or have appeared de novo as genomic innovations in streptophyte algae (Fig. 7G and fig. S6, A and B). Note that DTD2 was reported to be one of the only two proteins unique to Archaea and plants, while the other protein is topoisomerase VI subunit B (TopVIB) (20). Mutations in TopVIB are known to affect cell proliferation and endoreduplication, which are unique characteristics of the plant kingdom (47–50). While TopVIB is a subunit of archaean type VI topoisomerase, DTD2 is a stand-alone functional protein. Unlike DTD2, TopVIB is conserved in all the lineages of algae in addition to the land plants (fig. S6C). Therefore, this analysis indicates that DTD2 is the only gene unique to Streptophyta and Archaea.

**Recruitment of DTD2 from methanogenic archaea**

A thorough phylogenetic analysis of all the known archaean and plant DTD2s highlighted that among plant DTD2s, DTD2s of streptophyte algae are closer to the archaeal ones (Fig. 6A and fig. S5, A and B).

![Fig. 8. aa-tRNAs are hypersensitive to acetaldehyde. (A and B) Radar charts showing that aa-tRNA (α-Tyr-tRNA<sup>Tyr</sup>) and its analogs (α-Tyr2AA and Val3AA) are hypersensitive to acetaldehyde than other biomolecules (each corner represents the ratio of substrate to acetaldehyde, and hexagonal rings depict the percentage of the extent of modification. Related MS data are shown in (E) and figs. S7 and S8D. (C) TLC is showing differential migration of α-Tyr-tRNA<sup>Tyr</sup> at the different substrate-to-acetaldehyde ratios. aa-tRNA corresponding spot was observed at 1:0; monoethyl-aa-tRNA corresponding spots were observed at 1:1, 1:5, 1:10, and 1:100; and complete diethyl-aa-tRNA corresponding spot was observed at 1:1000 substrate-to-acetaldehyde ratios [related MS data are shown in (E)]. (D) TLC is showing differences in the spot positions of aa-AMP corresponding to the unmodified, monoethyl, and diethyl-Tyr-tRNA<sup>Tyr</sup> [lanes include untreated aa-tRNA, aa-tRNA incubated with only acetaldehyde, aa-tRNA incubated with only reducing agent, aa-tRNA incubated with both acetaldehyde (1:5 substrate-to-acetaldehyde ratio) and reducing agent, and aa-tRNA incubated with both acetaldehyde (1:1000 substrate-to-acetaldehyde ratio and it is indicated as asterisk in the figure) and reducing agent, respectively]. (E) MS data related to (C) (corresponding MS2 data are shown in fig. S8A).
Among archaeal DTD2s, *Methanocella conradii* shares maximum sequence identity with most of the DTD2s of streptophyte algae and land plants (Fig. 7, H and I). This emphasizes on a single acquisition event of DTD2 from methanogenic archaea to the plant kingdom that happened at the basal radiation of land plants. As mentioned earlier, except streptophyte algae, DTD2 is absent in all the other lineages of algae. This further strengthens that DTD2 was likely acquired from Archaea by streptophyte algae rather than by an endosymbiotic event. Since it is unlikely to be multiple gene loss events in the eukaryotic branch, we propose the likely possibility of a single acquisition event. In agreement, Methanocella is found to be in association with lower plants in bogs. Bogs are the oldest terrestrial ecosystems that are deprived of oxygen and enriched in d-amino acids (51, 52). One of the critical architectural innovations that happened during the evolution of land plants is the “root.” The root is the primary organ that is exposed to higher anaerobic stress and excess d-amino acids in the soil (2, 8, 13, 53, 54). In accordance, DTD2 is highly expressed in roots compared to the other organs (20). The confluence of the two independent stresses, namely d-amino acids and anaerobic stress, aggravates the formation of NEDATs. Therefore, the acquisition of DTD2 from Archaea in the terrestrial ecosystem by streptophyte algae likely benefited them to emerge as land plants during the Neoproterozoic Era (~800 million years ago).

### aa-tRNA is the first target to get modified by acetaldehyde

Chronic alcohol consumption in humans causes various cancers (21, 26, 27); the causative agent for the devastating effects is found to be acetaldehyde. Thus, acetaldehyde was demonstrated to be a genotoxic and teratogenic agent in mice and humans (21). Similar to aa-tRNAs, it mounts an electrophilic attack on the free amino groups of nucleic acids and proteins (22–25). To compare relative sensitivities of free amino acids, proteins, and nucleic acids with that of aa-tRNAs, each of these was treated with acetaldehyde. We found that aa-tRNAs are more prone to modification than other biomolecules. Amino acids and peptides are found to be the second preferred targets of acetaldehyde. The requirement of a 1000-fold molar excess or more of acetaldehyde to modify nucleotides suggests that nucleic acids are relatively less prone to modification. However, an equal amount of acetaldehyde is enough to modify aa-tRNAs (Fig. 8 and figs. S7 and S8, A to D). This clearly suggests that during anaerobic stress or acetaldehyde burst, aa-tRNAs are the first cellular components that undergo modification.

Since aa-tRNAs are the primary targets of acetaldehyde, DTD2 is expected to provide an early response during anaerobic stress. This is further supported by toxicity assays of DTD2 knockout plants that showed acetaldehyde sensitivity within a range of 0.01 to 1% (v/v) of acetaldehyde. Above the 1% (v/v) acetaldehyde, both wild-type and DTD2 knockout plants die (19, 20). This suggests that DTD2 operates to clear the toxic effects of low levels of acetaldehyde that are produced due to the physiological requirement of anaerobic fermentation. An excess acetaldehyde production leads to the modifications on other counterparts such as proteins and nucleic acids (Fig. 8, A, B, and E, and figs. S7 and S8D). In agreement, excess acetaldehyde not only generates ethyl modification on other biomolecules but also creates diethyl modification on aa-tRNAs. DTD2 does not deacetylate diethyl-d-aa-tRNAs (Fig. 8, C to E, and figs. S7, S8, A to C, and S9). The toxic effects of acetaldehyde are quenched by multiple tiers of detoxifying systems, especially ALDH and FANC2. ALDH, FANC2, and other repair mechanisms do not act directly on ethyl modification (21, 26, 27, 55–57), whereas DTD2 is the only enzyme that removes N-ethyl-d-amino acids directly from NEDATs (Fig. 9).

**DISCUSSION**

Our results show that acetaldehyde attacks a key component of translation, i.e., aa-tRNA, and creates ethyl modification. We also found that aa-tRNAs are hypersensitive to acetaldehyde compared to DNA and proteins. Acetaldehyde is a well-known causative agent of various cancers in alcohol-abused mice and humans (21, 26, 27). A major cause for such diseases is identified as ethyl adducts on DNA and proteins generated by acetaldehyde (22–25). In plants,
the anaerobic stress metabolite acetaldehyde is reported to accumulate during multiple environmental stresses that include floods, waterlogging, drought, cold, high salinity, and parasitic infections (3, 4, 8). Enhancing multistress tolerance in crop plants is a key priority in sustainable agriculture. It is therefore worth exploring the possibility of overexpression of DTD2 as a strategy for improving multistress tolerance in crop plants.

Acetaldehyde-modified aa-tRNAs (NEATs) are not acted upon by DTD1 and PTH. Except for plants, most eukaryotes and bacteria do not encode acetaldehyde biosynthesis machinery; hence, the presence of DTD2 is superfluous and is substituted by DTD1 to perform chiral proofreading in these organisms. Distribution of DTD2 is exclusive to Archaea and Streptophyta (including streptophyte algae and land plants), where they encounter the dual problem of acetaldehyde biosynthesis and α-amino acid enrichment. DTD2 and PTH share a common ancestral fold; moreover, PTH is the only known enzyme that acts on N-blocked aa-tRNAs (16). We speculate that during the evolution of Archaea, DTD2 evolved to circumvent early harsh geochemical conditions by stitching an additional domain to the existing basic scaffold of PTH. In addition, it is known that under extreme environmental conditions such as higher temperature, racemization rates of aa-tRNAs are higher and eventually could contribute to the accumulation of ϕ-aa-tRNAs to pernicious levels (58). Notably, PTH does not act on NELATs, which differ subtly with acetyl substrates and therefore delineates PTH’s specificity toward the carbonyl group that is attached to the α-amino group of amino acid. In contrast, DTD2 acts on ethyl modification but not on acetyl that clearly displays its specificity toward methylene group. Mechanistic insights into substrate specificity of DTD2 and PTH may provide the structural basis for their exclusive biochemical selectivity.

Land plant evolution is a major leap in the history of life that occurred during the Neoproterozoic era (~800 million years ago). Terrestrial flora played a pivotal role in changing the global environment and in the diversification of fauna on the land surface (59). Streptophyte algae successfully colonized the land with biochemical, cytological, developmental, and architectural adaptations (42–46). Investigations of the adaptations that occurred in streptophyte algae to be a successful colonizer of land are of extreme importance (42–46). In this regard, our finding of DTD2’s unique role in resolving anaerobic stress exacerbated by ϕ-amino acids brings another dimension to our understanding of the cellular and molecular events that led to the emergence of land plants. Moreover, this also provides the rationale for DTD2’s absence in Ophiokokona, which do not encounter sustained acetaldehyde and ϕ-amino acid stress (40, 41, 52). Notably, DTD2 is the only gene unique to Archaea and land plants and is the only known gene transfer event from Archaea to streptophyte algae. Moreover, the conservation of DTD2 from early-diverging streptophyte algae suggests that DTD2’s recruitment is an early event that likely facilitated the adaptation to terrestrial life (Fig. 9). Overall, our work sheds light on an unexpected cellular target of anaerobic stress and its mitigation, with implications for land plant evolution and multistress tolerance in crop plants.

MATERIALS AND METHODS
Materials
Materials were obtained from Merck unless otherwise mentioned. ESI-MS was performed using a Thermo Fisher Scientific Q-Exactive mass spectrometer. Plasmid pKPY514 coding for different subunits of E. coli phenylalanyl-tRNA synthetase (Ec PheRS) was a gift from D. Tirrell (Addgene plasmid #62598; http://n2t.net/addgene:62598; RRID: Addgene_62598) (60). Superdex 75 and sulfopropyl-Sepharose columns were purchased from GE Healthcare Life Sciences, USA. The nonhydrolyzable analogs of aa-tRNAs were purchased from Jena Biosciences, Germany.

Cloning, expression, and purification
Genes encoding DTD2s of P. horikoshii (Pho), M. jannaschii (Mj), and A. fulgidus (Af) and tyrosyl-tRNA synthetase (TyrRS) of T. thermophilus (Tth) were polymerase chain reaction–amplified from their genomic DNA using appropriate forward and reverse primers. A. thaliana (At) DTD2 gene was amplified from the complementary DNA of A. thaliana. Genes were cloned into the pET28b vector using restriction-free cloning (61). Primers used in this study are listed in table S1. Pho DTD2, Af DTD2, At DTD2, and Tth TyrRS were transformed and overexpressed into E. coli BL21(DE3), Ec PheRS into E. coli M15, and Mj DTD2 into BL21CodonPlus (DE3)-RIL strain of E. coli. Purification of 6× His-tagged proteins (Mj DTD2-C-His, At DTD2-N-His, Tth TyrRS-N-His, and Ec PheRS-N-His) was performed by Ni-NTA (nitrilotriacetic acid) affinity chromatography, followed by size exclusion chromatography (SEC). SEC was performed by using a Superdex 75 column (GE Healthcare Life Sciences, USA). Purification method and buffers for Ni-NTA and SEC were used as described (14). Untagged proteins (Pho DTD2 and Af DTD2) were purified using cation exchange chromatography (CEC) followed by SEC. For CEC, cells were sonicated in a buffer containing 50 mM bis-tris (pH 6.5) and 20 mM NaCl. The lysate was heated at 70°C for 30 min before subjecting to centrifugation (18,000 rpm for 30 min at 4°C). The supernatant was subjected to CEC column, and proteins were eluted with a gradient of NaCl from 50 to 200 mM. Sulfopropyl-Sepharose (GE Healthcare Life Sciences, USA) column was used for CEC. A buffer containing 100 mM tris (pH 8.0), 200 mM NaCl, 5 mM 2-mercaptoethanol (β-ME), and 50% glycerol was used to store all the purified DTD2 proteins.

Generation of ϕ-32P–labeled aa-tRNAs
All the tRNAs used in the study [E. coli (Ec) tRNA^Phe, P. horikoshii (Pho) tRNA^Phe, A. thaliana (At) tRNA^Phe, T. thermophilus (Tth) tRNA^Thr, and Mus musculus (Mm) tRNA^Thr (G4:U69)] were generated using the MEGAscript T7 Transcription Kit (Thermo Fisher Scientific, USA). tRNAs were end-labeled with ϕ-32P-ATP (adenosine 5′-triphosphate) (BRIT-Jonaki, India) using CCA-adding enzyme (62). Phenylalanylation of tRNA^Phe was done by incubating 1 μM tRNA^Phe with 2 μM E. coli PheRS in a buffer containing 100 mM Hepes (pH 7.5), 10 mM KCl, 30 mM MgCl2, 50 μM l-Phe or 50 μM l-Phe, and 2 mM ATP at 37°C for 15 min. Tyrosylation of tRNA^Phe, alanylation, serylization, and threonylation of tRNA^Thr (G4:U69) were done as mentioned earlier (14, 63). Aminoacylations were assessed and quantified as described below.

Generation of adducts on aa-tRNAs and their analogs
Adducts on aa-tRNAs and their analogs were generated by two different methods, namely single-step and two-step methods (fig. S8). Single-step method (probing relative acetaldehyde sensitivities of biomolecules; acetaldehyde titration with aa-tRNA)
A single-step method was used to find relative acetaldehyde sensitivities of aa-tRNAs (acetaldehyde titration with aa-tRNA), aa-tRNA
analogs, free amino acids, peptides, and nucleotides. In this method, 200 μM of each of aa-tRNA (Phe-tRNA^Phe^ and Tyr-tRNA^Tyr^), non-hydrolyzable analogs (d-Tyr2AA and l-Val3AA), amino acids (d-Tyr and l-Val), peptides (peptides 1 and 2), and nucleotides [adenosine 5’-monophosphate (AMP) and guanosine 5’-monophosphate (GMP)] were incubated with different concentrations of acetaldehyde [200 μM, 1 mM, 2 mM, 20 mM, 200 mM, and 2 M (stocks were prepared in ethanol)] “along with 400 mM NaCNBH₃” [stock was prepared in 100 mM potassium acetate (pH 5.4)] at 37°C for 30 min. Except for aa-tRNA, all the other samples were subjected to Eppendorf 5305 Vacufuge plus Concentrator to concentrate the sample. Samples were characterized using MS without any further processing. The method for processing and quantification of modification on aa-tRNA is discussed in detail below.

**Two-step method (generation of substrates for deacylation and thin-layer chromatography)**

A two-step method was used to generate ethyl modification on the aa-tRNAs that were used for deacylation assays. The final concentration of radiolabeled substrates (modified and unmodified aa-tRNAs) required for deacylation assays was 200 nM. Performing modification reactions with aa-tRNAs at such a low concentration of acetaldehyde (200 nM) was problematic because of its highly volatile nature. On the other hand, raising aa-tRNA concentration to 200 μM (concentrations used in single-step method) or more for each radiolabeled substrate that we used for biochemical assays is resource-intensive, and a large part of it will be unused because the final substrate concentration used for deacylation assays is only 200 nM. On the basis of these considerations, we used a two-step method to generate modification on aa-tRNA that gives maximum modification consistently. Substrates with homogeneous and maximum modification are a prerequisite to perform deacylation assays.

In the two-step method, acetaldehyde and reducing agents were added to aa-tRNAs or aa-tRNA analogs in two discrete steps. In the first step, 2 μM aa-tRNAs or 100 μM aa-tRNA analogs (used as a control) were incubated with 20 mM acetaldehyde at 37°C for 30 min. Before adding the reducing agent, the samples were subjected to Eppendorf 5305 Vacufuge plus Concentrator to “remove excess acetaldehyde” from the reaction. In the second step, the above “mixture was reduced with 400 mM NaCNBH₃” by further incubating at 37°C for 30 min. All the reactions were performed on a shaking incubator operated at 300 rpm. Acetaldehyde-treated and untreated analogs were characterized using MS without any further processing. In contrast, acetaldehyde-treated and untreated aa-tRNAs were ethanol-precipitated by overnight incubation at −30°C to remove salts and metal ions. After ethanol precipitation, pellets were resuspended in 5 mM sodium acetate (pH 5.4), and the same pellets were used for deacylation assays.

To check the aminoaetylation and modification status, reaction mixtures were analyzed before the ethanol precipitation step. One microliter of the reaction mixtures of modified aa-tRNAs or unmodified aa-tRNAs (control) was mixed with 2.5 μl of S1 nuclease (2 U/μl) (Thermo Fisher Scientific, USA) and incubated for 30 min at 22°C. One microliter of S1 nuclease digested samples was spotted onto cellulose F TLC plates (Merck KGaA, Germany). These TLC plates were developed using a mobile phase consisting of 100 mM ammonium chloride and 5% glacial acetic acid; the mobile phase front was allowed to move up to about three-fourth of the vertical length of the TLC plate before air-drying the plate. The developed TLC plates were exposed overnight to imaging plates (Fujifilm, Japan). Typhoon FLA 9000 biomolecular imager (GE Healthcare) was used for phosphorimaging of the exposed image plates. S1 nuclease is an endonuclease that hydrolyzes single-stranded RNA into mononucleotides. S1 digestion of tRNA therefore results in hydrolysis of single-stranded regions such as loops and the 3’-CCA end to free nucleotides. AMP, aa-AMP, and modified aa-AMP were visualized on TLC after phosphorimaging because the 3’-terminal nucleotide A76 was labeled with α-32P. Hence, AMP, aa-AMP, and modified aa-AMP that appear on TLC correspond to free/deacylated tRNA, aa-tRNA, and modified aa-tRNA, respectively (62). Percentages of aminoaetylation and modification were assessed using Image Gauge V4.0 software. Protocol for acetaldehyde modification assay was generated and optimized from the insights obtained from the earlier studies (24, 64). N-acetyl-aa-tRNAs were generated and quantified as described (31).

**Deacylation assays**

Deacylation experiments were performed by incubating different enzymes (PTHs, DTD1s, and DTD2s) of varying concentrations (as mentioned in the figures) with 0.2 μM of different substrates (α-32P-labeled modified or unmodified aa-tRNAs) in a buffer containing 20 mM tris (pH 7.2), 5 mM MgCl₂, 5 mM dithiothreitol (DTT), and bovine serum albumin (0.2 mg/ml) at 37°C. One microliter of the reaction mixture was withdrawn at different time points and subjected to S1 digestion (the remaining steps are the same to those discussed in the “Generation of adducts on aa-tRNAs and their analogs” section).

The percentage of aa-AMP or modified aa-AMP present at 0-min time point was considered as 100%. The percentage of aa-AMP or modified aa-AMP remaining after deacylation was plotted against the respective deacylation time points.

**Alkali treatment**

Aa-tRNAs, acetaldehyde-modified aa-tRNAs, and N-acetyl-aa-tRNAs were initially digested with S1 nuclease as mentioned earlier. S1-digested samples were later subjected to alkali treatment because alkaline conditions are incompatible with S1 digestion. Alkali treatment was carried out by incubating 100 nM S1-digested sample with 100 mM tris (pH 9.0) at 37°C. One microliter of the alkali-treated samples was directly spotted on to a TLC plate at different time intervals, and the latter steps were followed as described above (in the “Generation of adducts on aa-tRNAs and their analogs” section).

Half-life values were determined by using GraphPad Prism software by fitting the data points on to the curve according to the first-order exponential decay equation [S_t] = [S_0]e^{-kt}, in which the substrate concentration at time t is denoted as [S_t], [S_0] is the concentration of the substrate at time 0, and k is the first-order decay constant.

**Generation and characterization of NEDATs in E. coli**

*E. coli* K12Δtdt::kan cells were used to establish a surrogate system. Initially, primary cultures were grown at 37°C in 1x minimal salts with 0.2% maltose and kanamycin (50 μg/ml) (minimal medium). One percent of overnight-grown primary culture was inoculated to grow 400-ml secondary cultures (minimal medium with or without 3 mM d-tyrosine). Cultures were grown until OD₆₅₀ (optical density at 650 nm) reached to 0.8 [it was shown that at 0.8 OD₆₅₀, of the total Tyr-tRNA^Tyr^ pool, 40% was d-tyrosylated in the case of d-tyrosine–treated *E. coli* Δddt cells (65)]. Two hundred milliliters of each secondary culture was used for RNA isolation, and all the samples were...
treated with 0.5 mM NaCNBH₃ for 30 min at 37°C before RNA isolation. In the case of acetaldehyde-treated samples, before RNA isolation, cultures were additionally treated with either 0.005% (v/v) or 0.01% (v/v) acetaldehyde along with the reducing agent at 37°C for 30 min. Alkali treatment [5 mM CuSO₄ + 50 mM tris (pH 8.0)] was used to find out the formation of NEATs in acetaldehyde-treated cells (AA-tRNAs and modified aa-tRNAs co-migrate on the acrylamide gel electrophoresis (PAGE)). Hence, CuSO₄ + tris (8.0) treatment was carried out to distinguish modified aa-tRNAs from aa-tRNAs. Similar to alkaline treatment [100 mM tris (pH 9.0)], only aa-tRNAs are readily deacylated but not modified aa-tRNAs with CuSO₄ + tris (8.0) treatment. Retention (no hydrolysis) was only observed when cultures were grown in the presence of both D-amino acids and acetaldehyde that suggests accumulation of NEATs. A similar method was used in the case of N-acetyl-aa-tRNAs and peptideyl-tRNAs (66). Deacylations with both plant and archaeal DTD2s (similar to in vitro deacylation assays) were done to characterize the NEATs. RNA (0.25 absorbance at 260 nm unit) isolated from the different samples was subjected to alkaline treatment or deacylation with DTD2. After deacylation, RNA samples were analyzed by Northern blotting, for which the samples were first fractionated on 6.5% acid-urea polyacrylamide gel (denaturing PAGE analysis) for 20 to 24 hours at 4°C (67). The gel was then subjected to electrophoretoblotting on Hybond® membrane at 16 V, 5 A for 40 min. A 5'–end labeled DNA primer complementary to the anticodon loop of tRNA<sub>Tyr</sub> was used as a hybridizing probe (probe sequence: GCCAGATTACAGTCTGCTC). Radioactive 5'–end labeling of the probe was done with γ<sup>32</sup>P-ATP (BRIT-Jonaki, India) using polynucleotide kinase (NEB) as a reaction as recommended by the manufacturer. The hybridized blot was exposed overnight to image plate followed by phosphorimaging and quantification as mentioned above (in the “Generation of adducts on aa-tRNAs and their analogs” section).

**EF-Tu protection assay**

EF-Tu from *T. thermophilus* was activated using a protocol, as mentioned earlier (34). Due to lower activation efficiency (10 to 15%), the effective concentration of activated EF-Tu was assumed to be around 200 to 300 nM in a reaction mix containing 2 μM EF-Tu (68). EF-Tu protection assays were performed at 37°C in a reaction mix containing 2 μM activated EF-Tu or 2 μM unactivated EF-Tu, 0.2 μM 1-aa-tRNA or δ-aa-tRNA ([<sup>32</sup>P]-labeled) in 100 mM Hepes (pH 7.2), 2.5 mM DTT with 2 mM acetaldehyde, and 20 mM NaCNBH₃. A total of 1.5 μl of the reaction mixture was taken out at various time points and immediately mixed with an equal volume of 250 mM l-cysteine at 37°C to quench the unreacted acetaldehyde. These samples were selected to S1 nuclease digestion, and the latter steps for TLC were followed as discussed above (in the “Generation of adducts on aa-tRNAs and their analogs” section).

**Mass spectrometry**

To characterize modification status on aa-tRNAs (titrated with acetaldehyde), modified and unmodified aa-tRNAs (1 mM) were resuspended in 30-μl diethyl pyrocarbonate–treated water and mixed with 60 μl of aqueous ammonia [25% of (v/v) NH₄OH] and incubated at 70°C for 18 hours. This treatment resulted in hydrolysis of modified and unmodified amino acids and nucleotides from modified and unmodified aa-tRNAs. Hydrolyzed samples were further concentrated with Eppendorf 5305 Vacufuge plus Concentra-
support the phylogenetic tree. MrBayes has run with default parameters (prset aamodelpr = mixed, mcnc nchains = 1 ngen = 300,000). Clades with a bootstrap value above 90% were well supported, whereas values more than 70% are moderately supported, and the clades with bootstrap values less than 50% were considered unresolved. Web server iTOL (http://itol.embl.de) was used to visualize the phylogenetic tree.

**Statistical analysis**

All the experiments were performed in triplicates (unless otherwise mentioned). The mean values were used to plot the graphs, and each error bar denotes the SD from the mean value of three independent observations.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/7/6/eabe8890/DC1

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