

An amino-terminal point mutation increases EAAT2 anion currents without affecting glutamate transport rates

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Excitatory amino acid transporters (EAATs) are prototypical dual function proteins that function as coupled glutamate/ Na^+ / H^+ / K^+ transporters and as anion-selective channels. Both transport functions are intimately intertwined at the structural level: Secondary active glutamate transport is based on elevator-like movements of the mobile transport domain across the membrane, and the lateral movement of this domain results in anion channel opening. This particular anion channel gating mechanism predicts the existence of mutant transporters with changed anion channel properties, but without alteration in glutamate transport. We here report that the L46P mutation in the human EAAT2 transporter fulfills this prediction. L46 is a pore-forming residue of the EAAT2 anion channels at the cytoplasmic entrance into the ion conduction pathway. In whole-cell patch clamp recordings, we observed larger macroscopic anion current amplitudes for L46P than for WT EAAT2. Rapid L-glutamate application under forward transport conditions demonstrated that L46P does not reduce the transport rate of individual transporters. In contrast, changes in selectivity made gluconate permeant in L46P EAAT2, and nonstationary noise analysis revealed slightly increased unitary current amplitudes in mutant EAAT2 anion channels. We used unitary current amplitudes and individual transport rates to quantify absolute open probabilities of EAAT2 anion channels from ratios of anion currents by glutamate uptake currents. This analysis revealed up to 7-fold increased absolute open probability of L46P EAAT2 anion channels. Our results reveal an important determinant of the diameter of EAAT2 anion pore and demonstrate the existence of anion channel gating processes outside the EAAT uptake cycle.

After release from presynaptic nerve terminals, glutamate is quickly removed from the synaptic cleft by a family of glutamate transporters, the excitatory amino acid transporters (EAATs) (1–3). EAATs are not only secondary active glutamate transporters, but also function as anion-selective channels (4–7). EAAT anion channels regulate neuronal excitability and synaptic transmission (8, 9) as well as intracellular $[\text{Cl}^-]_{\text{int}}$ in glial cells (10). The importance of EAAT anion channels for normal cell function is emphasized by naturally occurring mutations that cause episodic ataxia and epilepsy and alter EAAT anion currents (11–15).

Glutamate transport is based on the translocation of the transport domain that encompasses binding sites for all transporter substrates. After association of glutamate, this domain moves across the membrane via a piston-like movement and then releases substrates to the intracellular membrane side. This so-called elevator mechanism was first described for the prokaryotic glutamate transporter Glt_{Ph} (16, 17) and later shown to also apply to an increasing number of other secondary active transporters (18–20). For EAAT/ Glt_{Ph} , it ensures strict stoichiometric coupling of glutamate to Na^+ , K^+ , and H^+ transport by permitting translocation only for certain ligation states of the transporter (21–24). Elevator-like transport is also the basis of the anion channel function of this class of glutamate transporters. In atomistic molecular dynamics simulations, the lateral movement of the transport domain and subsequent water entry in the cleft between transport and trimerization domain generates a selective anion conduction pathway (25). This novel conformation correctly predicts experimentally determined selectivities among anions and unitary anion currents and accounts for all published mutagenesis results on EAAT anion channels (25–29).

This mechanism of EAAT/ Glt_{Ph} anion channel opening permits the evolutionary optimization of transport rates without changes in anion currents and alterations of anion channel opening without modification in transport rates. It thus accounts for the existence of specialized glutamate transporters and low-capacity transporters with predominant anion channel function (4, 5, 30). To further test the predictions of this model, we searched for point mutations that modify anion channel open probabilities without altering glutamate transport rate. In a series of experiments in which EAAT residues that are homologous to anion pore-forming residues in Glt_{Ph} (25) were substituted by proline (because of its specific conformational rigidity), we found that L46P substantially increases macroscopic anion current amplitudes of EAAT2. We did not observe changes in the voltage and substrate dependence of mutant anion currents, suggesting that the glutamate transport cycle remained unaffected by this mutation. We here performed a detailed investigation of the consequences of this mutation on glutamate transport and anion conduction of EAAT2.

Results

L46P increases macroscopic EAAT2 anion currents

Fig. 1A depicts an alignment of amino-terminal sequences of mammalian SLC1 transporters and prokaryotic homologues. Bacterial transporters usually exhibit shorter amino-terminal

This article contains supporting information.

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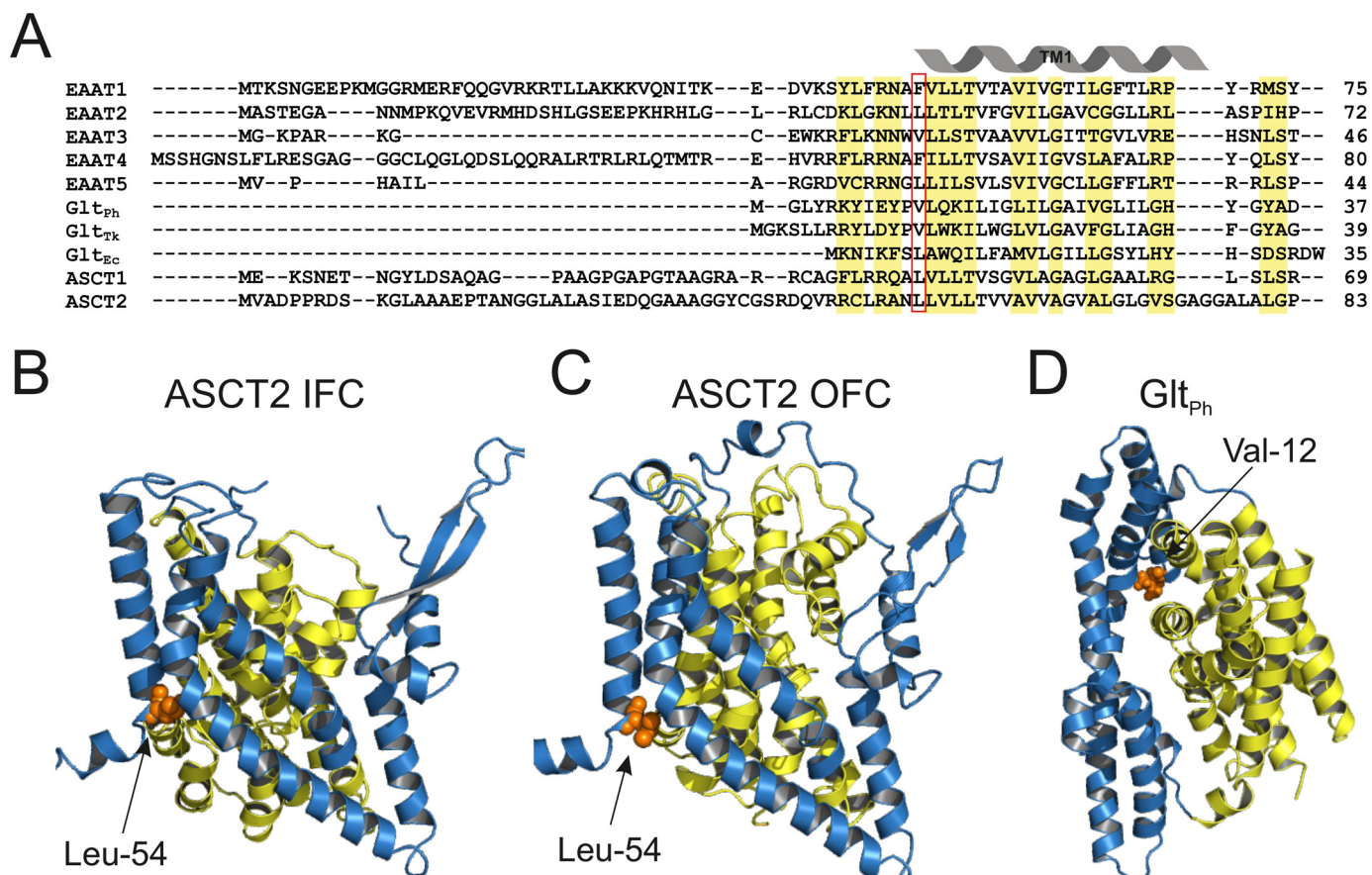


Figure 1. Leu-46 is a pore-forming residue and part of the trimerization domain. *A*, alignment of the amino terminus of EAATs, ASCTs, and prokaryotic homologs. Highly conserved regions are marked in yellow. *B–D*, position of the Leu-46-homologous residue Leu-54 in the ASCT2 topology model in the inward (32, 33) (*B*) and outward (31) (*C*) and of Val-12 in the Glt_{ph} open channel conformation (*D*) (25) (blue, trimerization domain; yellow, transport domain).

domains, and leucine 46 is not conserved among the family. At present, no three-dimensional structure exists for EAAT2, and we therefore mapped the homologous residue, leucine 54, to outward- (31) and inward-facing conformation (32, 33) of ASCT2, the only mammalian SLC1 transporter with known structures in both conformations (Fig. 1, *B* and *C*). Leucine 54 is close to the amino-terminal end of the trimerization domain and does not undergo movements during the isomerization from outward- to inward-facing conformations. The open channel conformation has so far not been described for ASCT2, and we therefore mapped the homologous valine 12 to Glt_{ph} in its open anion channel conformation (25), where it projects its side chain into the water-filled anion permeation pathway as pore-lining residue (25) (Fig. 1*D*).

Fig. 2, *A* and *B* depict representative recordings from cells transiently transfected with WT (Fig. 2*A*) or L46P EAAT2 (Fig. 2*B*). In these experiments, cells were dialyzed with KNO₃-based internal solution and then perfused either with NaNO₃-based external solutions without or with 1.0 mM L-glutamate, or with an external solution in which NaNO₃ was completely substituted with KNO₃. EAAT2 transports glutamate in the presence of external Na⁺ and L-glutamate (21–23). The use of NO₃[−] that is more permeant than Cl[−] (34) increases EAAT2 anion currents to levels greatly exceeding uptake currents (5, 35) (Fig. 2, *C* and *D*). Almost complete block upon application of TBOA, which prevents opening of EAAT anion channels by blocking

inward movement of the transport domain (36), demonstrates that WT and mutant EAAT2 anion currents are significantly larger than background currents (Fig. 2, *C* and *D*) and can thus be directly measured without subtraction procedures for many experimental conditions (35, 37).

WT as well as L46P EAAT2 anion currents are small and without appreciable time and voltage dependence in NaNO₃-based external solutions lacking glutamate (Fig. 2). Application of L-glutamate increases anion currents for WT and mutant transporters (Fig. 2, *C* and *D*). EAAT2 also mediates large anion current amplitudes under conditions, in which extracellular Na⁺ is completely substituted by K⁺ (38, 39). WT and mutant currents displayed similar time, voltage, and substrate dependences. However, L46P EAAT2 anion currents are much larger than WT currents. Moreover, whereas WT EAAT2 anion current amplitudes were highest in external Na⁺ and L-glutamate, we observed maximum mutant current under KNO₃-based external solutions (Fig. 2*D*).

L46P affects neither expression nor subcellular distribution of EAAT2

To test whether increased macroscopic L46P EAAT2 currents arise from changes in subcellular localization or in expression levels, we employed confocal imaging and protein biochemistry (Fig. S1). WT and L46P EAAT2 display predominant surface membrane insertion in confocal images from HEK293T cells

A mutation increasing EAAT anion channel open probability

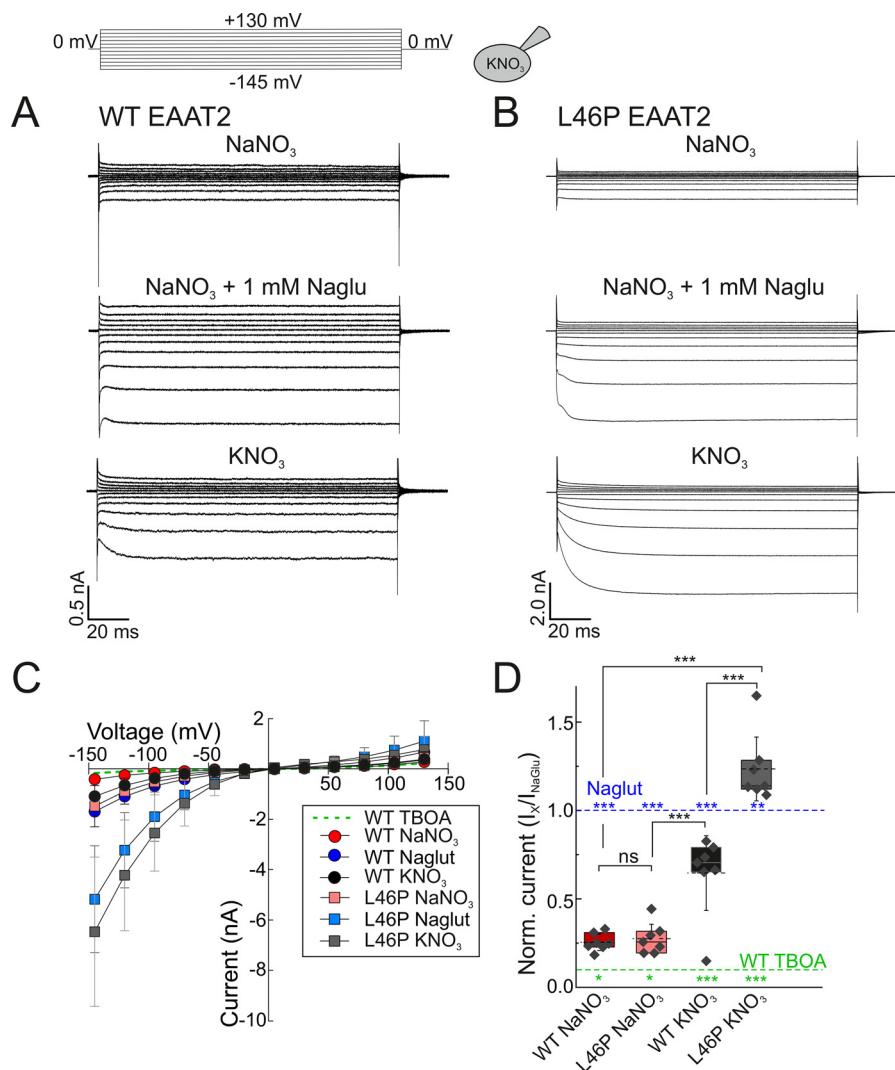


Figure 2. L46P increases EAAT2 anion currents under multiple ionic conditions. A and B, representative current recordings from HEK293T cells expressing WT (A) or L46P (B) EAAT2. Cells were intracellularly dialyzed with a KNO_3 -based solution and perfused with NaNO_3 -based solution without or with glutamate, or with KNO_3 -based solutions. C, voltage-dependence of mean steady-state currents from cells expressing WT or L46P EAAT2. The background current (dotted green line) was obtained from cells expressing WT EAAT2 perfused with TBOA. D, normalized current amplitudes at -145 mV from cells expressing WT or L46P EAAT2. All error bars represent 95% CI with $n = 7$. To account for larger L46P EAAT anion current amplitudes, WT and mutant currents are shown on different scales. Data are significant different with *, $p \leq 0.05$; **, $p \leq 0.01$; ***, $p \leq 0.001$.

(Fig. S1, A and B). Fig. S1C depicts fluorescent scans of SDS-PAGE from membrane preparation of transfected HEK293T cells, indicating core and complex glycosylated WT and L46P EAAT2 (40). Individual bands were quantified with ImageJ software and revealed similar expression levels (Fig. S1C). To exclude the possibility that changes in the number of transfected cells compensate for altered expression levels, we also determined transfection rates by manually counting transfected and untransfected cells. We did not observe differences between WT and mutant transporters (Fig. S1D) and conclude that L46P leaves expression levels and subcellular distributions of EAAT2 unaffected.

L46P slightly increases unitary current amplitudes of EAAT2 anion channels

Because expression levels of WT and mutant transporters are similar, the observed differences in L46P EAAT2 macro-

scopic current amplitude might be because of increased mutant anion channel conductances or open probabilities. We employed nonstationary noise analysis to determine unitary current amplitudes of WT and L46P EAAT2 anion channels (41). To perform such experiments with comparable WT and mutant current amplitudes, we generated inducible stable Flp-In T-Rex 293 cell lines for WT as well as for L46P EAAT2 (40).

In WT cells, anion currents with amplitudes sufficient for noise analysis were only rarely observed, even under maximum tetracycline concentration and induction periods. Flp-In T-Rex 293 cells differ from HEK293T in stable expression of the lacZ-Zeocin fusion gene and the Tet repressor, modifications that are not expected to modify the function of EAAT2, making the comparison of noise analysis results from both types of cells possible. We tested potential effects of the expression system by comparing noise analysis results on WT EAAT2 anion currents in HEK293T and Flp-In T-Rex 293 cells (Fig. S2). Both

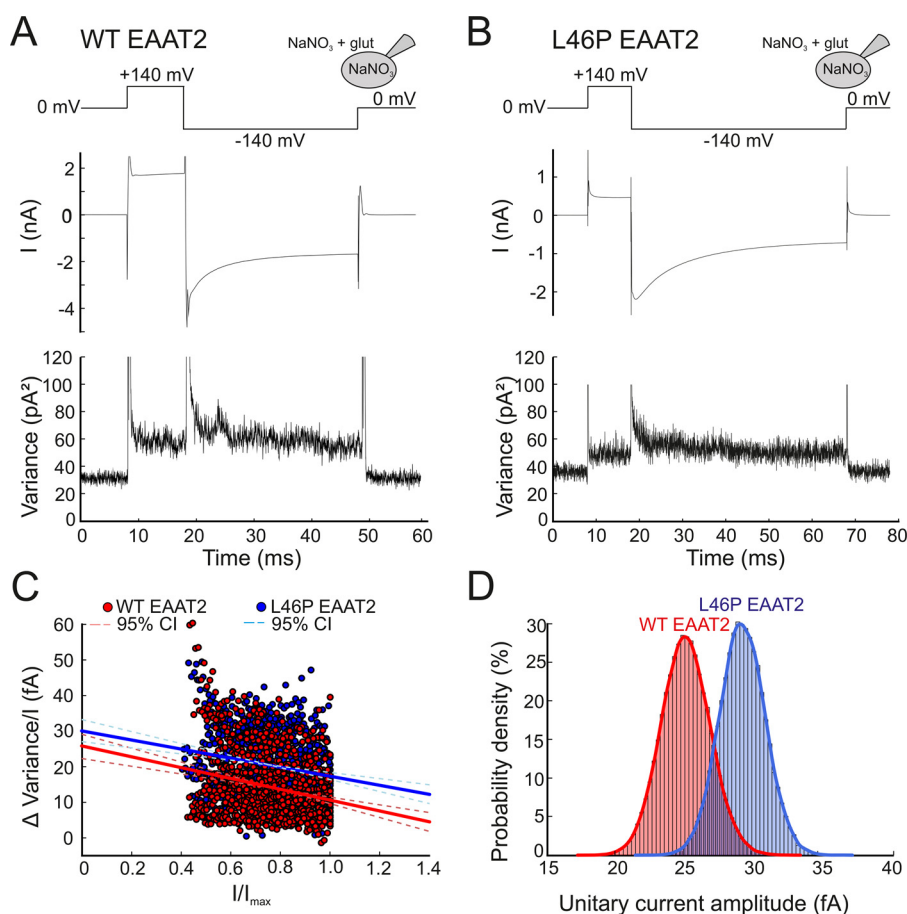


Figure 3. L46P slightly increases unitary current amplitudes. A and B, voltage protocol and time courses of representative current amplitudes and current variances from HEK293T cells transiently transfected with WT (A) or Flp-In T-Rex 293 cells stably expressing L46P (B) EAAT2 measured in symmetrical NaNO_3 based solutions. C, pooled noise analysis data from 20 cells expressing WT (red circles) and 17 cells expressing L46P EAAT2 (blue circles). For comparison between different cells, current variance by mean current amplitude ratios are plotted against normalized current amplitudes. Solid lines represent fits with linear functions and dashed lines give the 95% confidence interval of the fit determined by bootstrap regression analysis. D, distribution of estimated unitary current amplitudes for WT and L46P EAAT2 derived from 50,000 bootstrap samples of the original data. Histograms have been normalized such that the integral over the range is 1 and fitted with a Gaussian function (WT EAAT2, $\mu=24.8$, $\sigma=1.75$; L46P EAAT2, $\mu=28.9$, $\sigma=1.64$).

systems provided the same results, and we pooled results from both expression systems for the analysis of WT EAAT2. For L46P EAAT2, noise analysis was exclusively performed in inducible Flp-In T-Rex 293 cells.

Fig. 3, A and B show representative time courses of mean macroscopic currents and corresponding current variances for WT (A) or L46P (B) EAAT2 elicited by repetitive voltage steps to -140 mV in symmetrical NaNO_3 and 0.5 mM external L-glutamate. Fig. 3C provides pooled noise analysis data from 20 cells expressing WT (red circles) and 17 cells expressing L46P EAAT2 (blue circles). We plotted ratios of current variances by mean current amplitudes against normalized mean current amplitudes at various time points. A fit of the resulting relationship with a linear function (Equation 3) provides the single-channel current amplitudes as y axis intercept (42). Bootstrap analysis was used to simulate the error-generating process of data sampling from a larger population and to get an approximation of the true error of fitting parameters. 50,000 bootstrap samples (each containing 1000 WT or 850 mutant EAAT2 data points with replacements) were randomly generated from the original dataset (41, 43) and individually fitted with linear functions (Fig. 3C). Unitary current amplitudes were determined

for each bootstrap sample from linear fits, providing single channel amplitudes of 25 ± 2 femtoampere (fA) (mean $\pm$ 95% CI; $n=20$) for WT and 29 ± 2 fA (mean $\pm$ 95% CI; $n=17$) for mutant EAAT2 ($p < 0.05$) (Fig. 3D). We conclude that L46P slightly increases the unitary current amplitude of EAAT2 anion channels.

L46P affects pore dimensions of EAAT2 anion channels

EAAT anion channels conduct a variety of anions; they select, however, against large anions by size (25, 34, 35). The minimum pore size of the GltpH anion channel is around $5\text{--}6$ Å (25), only slightly smaller than the minimum size required for gluconate permeation (6.9 Å) (44). To test for potential changes in selectivity of L46P EAAT2 anion channels, we recorded currents in cells dialyzed with NaNO_3 and perfused with solutions containing mixtures of NaNO_3 and Na-gluconate or of NaNO_3 and choline- NO_3 (Fig. 4). In these experiments, accurate separation of EAAT2-mediated currents from endogenous current amplitudes is especially important. We therefore measured background currents by adding TBOA at the end of each experiment and corrected WT and mutant EAAT2 anion currents for such current components.

A mutation increasing EAAT anion channel open probability

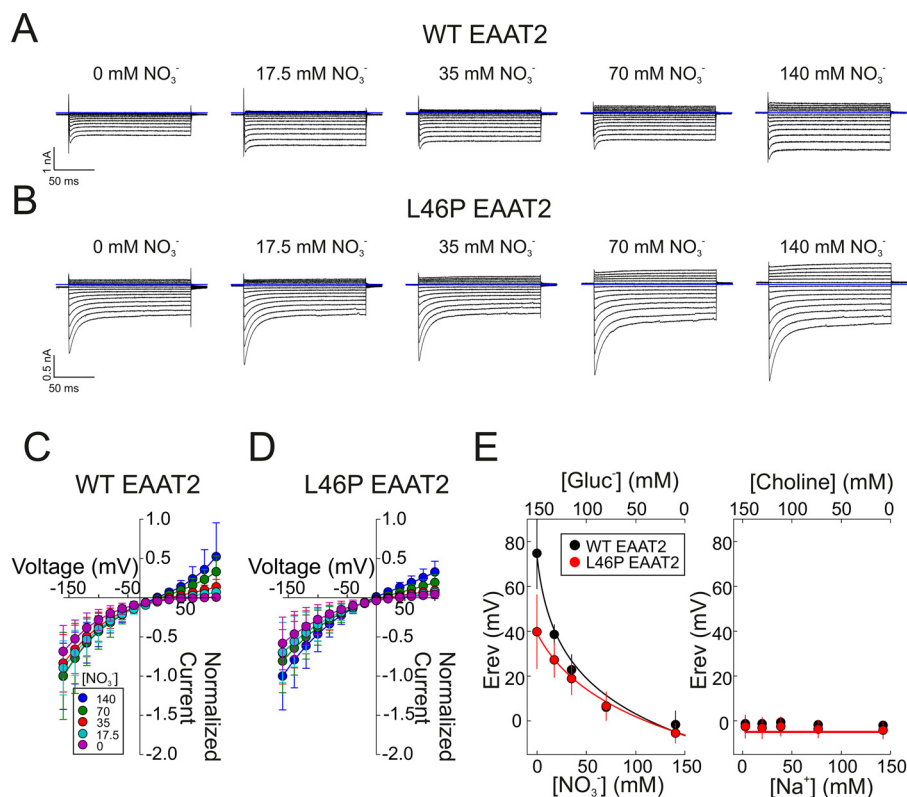


Figure 4. L46P EAAT2 anion channels are permeable to gluconate. A and B, representative anion current recordings from HEK293T cells either expressing WT (A) or L46P (B) EAAT2, subsequently perfused with 1 mM glutamate and different external $[\text{NO}_3^-]$ (substituted with gluconate) and internally dialyzed with a NaNO_3 -based solution. EAAT2 anion currents were corrected for background measured in glutamate-free solution supplemented with 0.1 mM DL-TBOA. C and D, normalized and background-corrected current-voltage relationships from cells expressing WT (C) or L46P (D) EAAT2 at various external $[\text{NO}_3^-]$. E, changes in WT or L46P EAAT2 anion current reversal potentials upon variation of external $[\text{NO}_3^-]$ or $[\text{Na}^+]$. Solid lines represent fits with the Goldman-Hodgkin-Katz equation. All error bars indicate mean $\pm$ S.D.

Outward currents were virtually nonexistent in cells expressing WT transporters perfused with NO_3^- -free external solutions (Fig. 4, A and C), in full agreement with a negligible gluconate $^-$ permeability. In contrast, we observed significant outward currents for L46P EAAT2 (Fig. 4, B and D). Gluconate $^-$ permeability of L46P EAAT2 suggests that the mutations widen the selectivity filter and permit passage of anions that are usually too large to permeate.

Plotting current reversal potentials *versus* external [gluconate $^-$] or external $[\text{Na}^+]$ (Fig. 4E) demonstrate that WT and mutant transporters differed significantly in their relative gluconate permeability, whereas neither WT nor L46P EAAT2 is measurably permeable to Na^+ . Permeability ratios obtained from the Goldman-Hodgkin-Katz (Equation 1) provided $P_{\text{Gluc}}/P_{\text{NO}_3}$ values of 0.18 ± 0.02 (mean $\pm$ S.D. from 50,000 bootstrap samples) for L46P, as compared with 0.05 ± 0.01 (mean $\pm$ S.D. from 50,000 bootstrap samples) in WT transporters.

Surprisingly, no L46P EAAT2 currents were observed in symmetric gluconate (see Fig. 6A). We thus postulate that NO_3^- binding may induce a transient widening of the L46P EAAT2 selectivity filter, which allows for subsequent gluconate $^-$ permeation.

WT and L46P exhibit similar unitary transport rates

WT and L46P EAAT2 differ neither in unitary anion currents nor in expression levels, making enhanced absolute open probabilities of mutant anion channels a likely reason for the

large anion currents in cells expressing L46P EAAT2. Recent work has demonstrated that nonstationary noise analysis cannot accurately determine absolute open probabilities of EAAT anion channels (12, 45). We therefore decided to estimate the fraction of time in which EAAT transporters assume the open anion channel conformation, using consecutive measurements of coupled transport and anion currents at the same cell (45). Because this analysis requires knowledge not only about unitary anion current amplitudes, but also of individual transport rates, we measured recovery rates of transport current depression using fast substrate application on excised outside-out patches to measure these values (46, 47).

Under conditions permitting forward transport, *i.e.* intracellular K^+ and extracellular Na^+ , fast application of L-glutamate initiates transport cycles and generates a transient capacitive current followed by a steady-state transport current (Fig. 5). The amplitudes of both current components are proportional to the number of transporters accessible to external glutamate. In a double pulse protocol with two subsequent glutamate applications, the amplitude of the second electrical signal depends on the time period between the two pulses. If the second pulse is applied immediately after the first pulse, the majority of transporters will assume intermediate positions of the transport cycle and only a few transporters will be accessible to external glutamate. With increasing duration between two subsequent glutamate applications, the current amplitude will

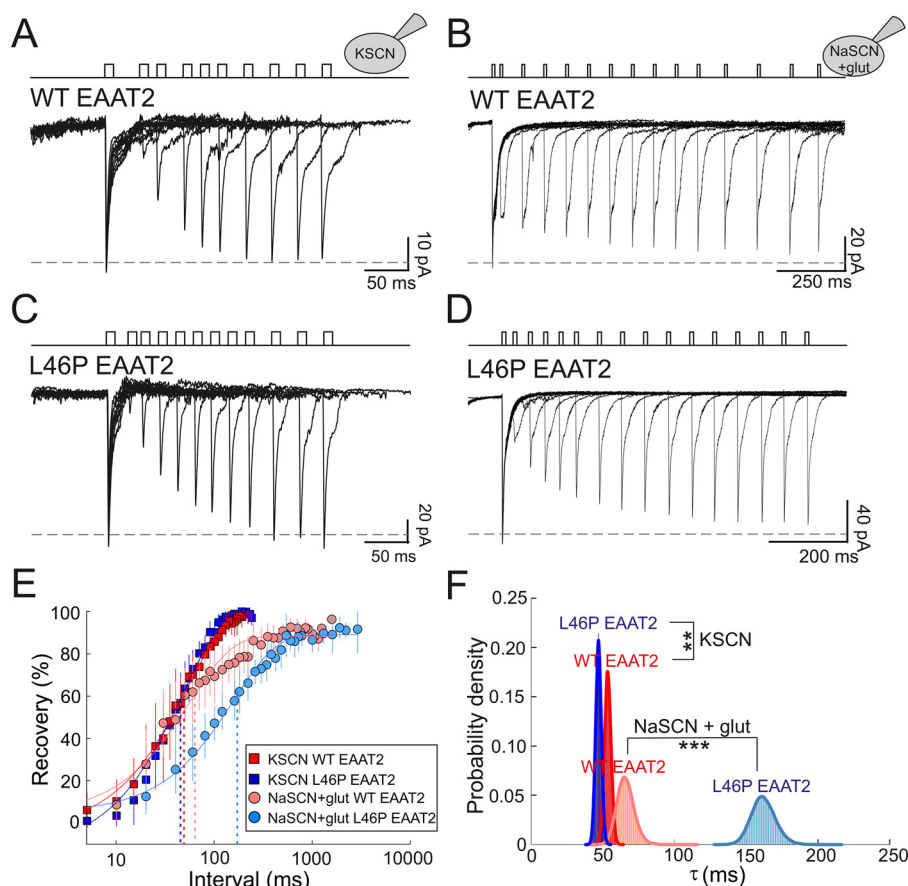


Figure 5. L46P leaves EAAT2 forward transport unaffected. A–D, representative rates of WT (A and B) or L46P (C and D) EAAT2 transporter current recovery from depression. Experiments were performed on excised outside-out patches either with KSCN- (A and C) or NaSCN/L-glutamate-based (B and D) cytoplasmic solutions. E, mean ratios of second response by first response peak current amplitudes plotted against the interval between pulses for 115 mM [KSCN]_{in} from patches containing WT (red squares, $\tau = 53.3 \pm 2.3$ ms, $n = 10$) or mutant EAAT2 (blue squares, $\tau = 46.9 \pm 2.0$ ms, $n = 8$), or for 115 mM NaSCN/10 mM L-glutamate-based solutions from patches containing WT (pink circle, $\tau = 65.9 \pm 6.0$ ms, $n = 8$) or mutant EAAT2 (light blue circle, $\tau = 161.9 \pm 8.4$ ms, $n = 6$). Error bars indicate mean $\pm$ S.D. F, probability density of transport rates obtained with 50,000 bootstrap samples randomly selected from the original dataset. Data are significant different with *, $p \leq 0.05$; **, $p \leq 0.01$; ***, $p \leq 0.001$.

recover with a time course that is determined by the average time required for a complete transport cycle (46, 47).

Fig. 5 depicts representative current responses to subsequent 10 ms application of 10 mM L-glutamate at increasing intervals for WT (Fig. 5, A and B) and L46P EAAT2 (Fig. 5, C and D). With K⁺-based pipette solutions, the recovery was fitted with a single exponential function with time constants of 53.3 ± 2.3 ms (mean $\pm$ S.D. from 50,000 bootstrap samples; $n = 10$) for WT and slightly faster values (46.9 ± 2.0 ms; mean $\pm$ S.D. from 50,000 bootstrap samples; $n = 8$; $p < 0.005$) for L46P EAAT2 (Fig. 5, E and F). We conclude that L46P causes only slight changes in rate-limiting steps of forward glutamate transport.

We next performed comparable experiments under homoexchange conditions, with intracellular solutions containing Na⁺ and L-glutamate (Fig. 5, B and D). Under these conditions, full transport cycles are impossible and recovery rates are determined by re-translocation and unbinding of L-glutamate. Under this condition L46P decelerated the time course of recovery (65.9 ± 6.0 ms (mean $\pm$ S.D. from 50,000 bootstrap samples; $n = 8$) for WT and 161.9 ± 8.4 ms (mean $\pm$ S.D. from 50,000 bootstrap samples; $n = 6$) for L46P EAAT2) (Fig. 5, E and F). Forward transport and homoexchange transport cycles

differ in the re-translocation step from inward- to outward-facing conformations. Because recovery time constants are significantly longer under homoexchange conditions than under forward transport, the re-translocation step must be the rate-limiting transition in both cases. Thus, L46P decelerates L-glutamate-bound re-translocation and/or substrate dissociation from the outward-facing conformation of EAAT2 transporters.

Absolute open probabilities of L46P EAAT2 anion channels are significantly increased

We next studied whole-cell currents in cells intracellularly dialyzed with a K-gluconate-based pipette upon subsequent perfusion with Na-gluconate-, NaNO₃-, or KNO₃-based external solutions (45) (Fig. 6). Electrogenic glutamate transport can be measured in isolation as glutamate-induced EAAT2 currents, when permeable anions are completely substituted with gluconate (5). Under these conditions, we observed comparable current amplitudes for WT and L46P EAAT2 at -100 mV (Fig. 6), in full agreement with unaltered expression and subcellular distribution (Fig. S1) and only slight differences in individual transport rates of mutant transporters (Fig. 5). However, application of L-glutamate-supplemented NaNO₃-based external

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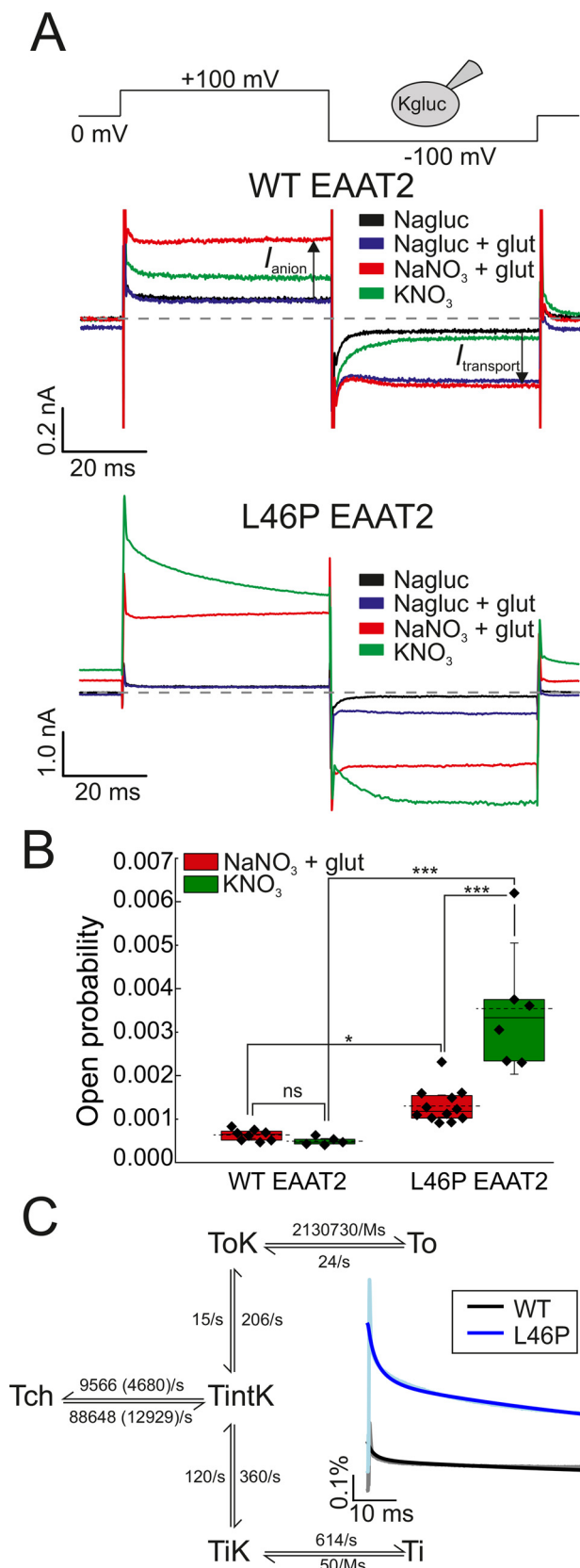


Figure 6. L46P increases absolute open probabilities of EAAT2 anion channels. A, representative current responses of HEK293T cells expressing WT or L46P EAAT2 to consecutive voltage steps to +100 mV and −100 mV. Cells were dialyzed with a K-gluconate-based pipette solution and consecutively perfused with Na-gluconate-based solution with or without L-glutamate, with a NaNO₃-based solution containing 0.5 mM L-glutamate, and

solution revealed dramatic difference in WT and mutant anion current amplitudes at positive potentials (+100 mV). Because of the pronounced voltage dependence of glutamate uptake currents (5), these currents are predominantly conducted by EAAT2 anion channels. Subsequent change to KNO₃-based external solutions resulted in current amplitudes that are only slightly smaller than for Na⁺/glutamate in WT and larger in L46P EAAT2 (Fig. 6, A and B).

To obtain the number of transporter subunits in the studied cells, we divided the transport current amplitude at −100 mV by individual transport rates of WT and L46P EAAT1 (Fig. 5). We next applied this number together with the unitary anion current amplitude obtained in Fig. 3 to calculate absolute open probabilities for WT and mutant EAAT2 (45) (Fig. 6B). We determined an absolute open probability of 0.06% ± 0.01% (mean ± 95% CI; *n* = 9) for WT and 0.13% ± 0.03% (mean ± 95% CI; *n* = 12) for L46P EAAT2 at +100 mV with external NaNO₃ and L-glutamate. L46P thus causes 2-fold larger open probability for EAAT2 anion currents than for WT under forward transport conditions and a 7-fold increase under K⁺-exchange conditions (0.05% ± 0.01% (mean ± 95% CI; *n* = 5) for WT and 0.35% ± 0.15% (mean ± 95% CI; *n* = 6) for L46P EAAT2 at +100 mV) with external KNO₃ (Fig. 6B). Because L46P EAAT2 does not reach a steady state current under K⁺-exchange conditions, an exponential function was fitted, and the instantaneous current 1 ms after pulse application was used to calculate the open probability.

External application of NaNO₃ or KNO₃ to cells expressing L46P EAAT2 did not only increase outward currents at positive potentials, but also inward currents at negative voltages (Fig. 6B). Such currents, which were absent in cells expressing WT EAAT2, are because of the gluconate permeability through L46P EAAT2 anion channel in the presence of NO₃[−] (Fig. 4).

With KNO₃ on both sides, voltage steps to positive potentials elicit prominent deactivating L46P EAAT2 anion currents, which were not observed in experiments with WT EAAT2 (Fig. 6A) or other mutants (24). Kinetic modeling demonstrates that the specific current kinetics are a direct consequence of the modification in anion channel opening/closing transitions in L46P EAAT2 (Fig. 6C). Under these conditions, the number of functional transporter states are significantly decreased, permitting direct quantification of particular reaction rates by fitting the time courses of anion channel open probabilities. Averaged time courses of absolute open probabilities in response to voltage steps to +100 mV for WT and L46P EAAT2 were generated by normalizing current responses to numbers of transporters per individual cell and mean unitary anion current amplitudes and fitted to a transport/anion channel scheme

finally with a KNO₃-based solution. B, estimated anion channel open probabilities for WT and mutant EAAT2 anion channels at Na⁺- or K⁺-based external solutions. Data are shown as mean ± 95% CI. Data are significant different with *, *p* ≤ 0.05; **, *p* ≤ 0.01; ***, *p* ≤ 0.001. C, absolute open probabilities, kinetic model, and simulated absolute open probabilities for WT and L46P in symmetric K⁺. Bright lines represent measured data, dark line represents modeled currents. Absolute open probabilities were calculated for five cells and averaged. Model rates that are different for L46P are given in parentheses.

(Fig. 6C) using a genetic algorithm (shown as *colored lines*). We assumed that anion channel (T_{Ch}) opening occurs from intermediate transport conformation ($T_{int}K$), which are not anion conducting, as predicted by recent simulation results (25). WT and L46P fit parameters only differ in anion channel opening and closing, indicating that L46P predominantly modifies EAAT2 anion channel gating.

Discussion

Channels and transporters mediate transmembrane ion transport employing fundamentally different mechanisms. Whereas ion channels support ion diffusion through aqueous conduction pathways, conformational changes are responsible for transmembrane ion translocation in transporters. This clear dichotomy has been challenged by the identification of channel modes in secondary active transporters, in which these proteins conduct currents via the transient formation of ion-selective pores (7). A comprehensive understanding of the molecular bases and the cellular functions of such channel modes is still lacking. Recent insights about the structural basis of their anion channel mode (25) and potential cellular functions (8–10) have made the EAAT glutamate transporter family a particularly well-understood example of dual function transport proteins.

Computational and experimental data support the notion that EAATs/Glts form anion-selective conduction pathways at the interface between trimerization and transport domain via lateral movement of the transport domain in intermediate conformations (25, 28, 29). Progression of the transport cycle requires anion channel closure (25), and efficient glutamate transport is only possible at the cost of low anion channel open probability. Functional analysis of *SLC1A3* mutations associated with episodic ataxia type 6 revealed that excessive EAAT anion currents might perturb brain development and function (11–14). The interdependency between glutamate transport and anion channel activity together with the pathological consequences of increased EAAT anion channel activity demonstrate the evolutionary need for a strict regulation of anion channel gating.

We found a mutation in EAAT2 that substantially increases macroscopic anion current amplitudes, without discernible effects in its voltage and substrate dependence and studied the functional consequences of this L46P mutation on glutamate transport and unitary properties of EAAT2 anion channels. We demonstrate that L46P leaves protein expression and subcellular distribution of EAAT2 unaffected (Fig. S1), but augments anion currents (Fig. 2). The increased anion current amplitudes are not because of changes in unitary current amplitudes (Fig. 3), but rather caused by an increased probability of occupying open anion channel conformations (Fig. 6). Moreover, L46P modifies EAAT2 anion channel selectivity, resulting in significant permeability of gluconate[−] (Fig. 4).

The dual function of EAATs as transporter and as anion channel, and the resulting complex gating mechanism underlying EAAT anion channel opening, makes the determination of absolute open probabilities difficult. For many ion channels, nonstationary noise analysis accurately quantifies unitary current amplitudes together with the number of channels in the

evaluated cell/membrane patch. However, this method does not provide reliable numbers for EAATs. Noise analysis on EAAT anion currents provided rather large open probabilities (35, 37), with values well above 0.5. Recently, a disease-causing point mutation was reported that decreases the number of transporters in the surface membrane but increases macroscopic anion current amplitudes more than 5-fold (12) at unaltered unitary current amplitudes. This observation demonstrates that absolute open probabilities of EAAT anion channels are not correctly determined using noise analysis. A possible reason for this inaccuracy has been already described for the anion-proton exchanger CLC-4 that also assumes anion channel modes in addition to functioning as secondary active transporter (42). Very fast transitions in combined transport/anion channel cycles cause semi-equilibria between transporters with open and closed channels. Hence, only transporters, which undergo rapid transitions between open and closed channel, will be counted by noise analysis, resulting in underestimation of the number of transporters and overestimation of absolute open probabilities (42).

We used an alternative approach that is based on the comparison of glutamate transport currents and anion currents (45). We first determined transport rates of WT and L46P EAAT2 (Fig. 5) and then used these transport rates to calculate the numbers of transporters in individual cells from measured glutamate uptake currents (Fig. 6). Dividing macroscopic anion current amplitudes by the corresponding transporter number and the unitary current amplitude (Fig. 3) provides absolute open probabilities (Fig. 6). This analysis reveals that WT EAAT2 spent only 0.06% of the total time in the anion conducting state under ionic conditions that permit forward transport, and an absolute open probability of 0.05% in the K^+ exchange mode. These values are very low and are fully consistent with EAAT2 being an effective glutamate transporter with rather low associated anion current (5). L46P increases this value to 0.13% in the forward mode and to 0.35% in the K^+ exchange mode.

Our analysis of L46P EAAT2 illustrates that changes in anion channel open probability can occur without concomitant alterations in glutamate uptake. We determined individual transport rates by measuring recovery rates of transporter current depression in the forward transport mode (Fig. 4) and observed closely similar values for WT and L46P EAAT2. L46P thus does not cause major changes in forward transport rates. However, L46P EAAT2 anion currents, either in absolute values (Fig. 2) or after normalization to transport rates (Fig. 6), are much larger. We fitted a kinetic scheme, developed to describe secondary active glutamate transport (46) and refined to describe anion channel opening from intermediate conformations to WT and L46P EAAT2 anion currents, for one ionic condition, K^+ on both sides. We found that sole modification of transitions between intermediate transport conformation and open channel conformation fully describes L46P-mediated changes in EAAT2 anion channel amplitudes and kinetics (Fig. 6C).

Our estimates of glutamate transport rates resemble data with native neuronal EAATs in hippocampal Purkinje neurons (48) but are smaller than published values on heterologously expressed (49) or native (46) EAAT2. At present, we do not

know the origin for these differences. They might be because of small variation in the composition of the internal and external solutions. Otis and Kavanaugh (49) use higher internal $[K^+]$ and apply TEA to the cytoplasmic membrane, rather than to the external solution as we did in our experiments. There are earlier reports on variation of recovery rates depending on cell systems and experimental approach (47). L46P has a slightly more pronounced effect on anion current amplitudes in the K^+ exchange mode than under glutamate forward transport conditions (Figs. 2 and 6). This might be because of a K^+ dependence in anion channel opening. Alternatively, they might indicate subtle alterations of certain transport transitions. L46P has only minor effects on recovery rate of depression of the transporter current under forward transport conditions (Fig. 5), indicating similar glutamate transport rates for WT and mutant transporters. However, under Na^+ -glutamate exchange conditions, recovery time constants of L46P EAAT2 are more than 2-fold larger than for WT (Fig. 5). These differences suggest that L46P additionally affects Na^+ /glutamate-bound retranslocation and/or release. Neither in an inward-facing ASCT2 structure nor in the EAAT1 structure, is L46P directly interacting with the transport domain. However, it is in close proximity to the gate to the substrate binding sites (HP2) so that local conformational changes could translate in different translocation or glutamate release kinetics.

Earlier work comparing glutamate transport and anion current amplitudes permitted the distinction of EAAT isoforms that mainly operate as glutamate uptake carriers (EAAT1, EAAT2, EAAT3) (5) and others, EAAT4 and EAAT5, that exhibit low glutamate transport rates and predominantly function as glutamate-gated anion channels (4, 50). Published data on EAAT1 and EAAT3 depict smaller anion than uptake currents (5), indicating very small anion channel open probabilities also for these two isoforms. Such experiments have not been possible for EAAT4 or EAAT5 because of the low glutamate transport rates of these two isoforms (30, 51). However, the recent comparison of a mouse EAAT2 splice variant, GLT-1c, and mouse EAAT5 in the same expression system with identical experimental approaches for both isoforms (41) revealed similar macroscopic anion current amplitudes at comparable expression levels and subcellular distribution. This finding argues against major differences in absolute open probability between high- and low-capacity glutamate transporters. EAAT4 and EAAT5 have been shown to exhibit very small transport rates, suggesting that the main difference between specialized EAAT transporters and EAAT anion channels is the transport rate rather than the anion channel open probability (30, 51). However, the small open probability of EAAT anion channels permits adjustment of open probabilities without changes in transport rates.

In summary, we here present a point mutation that exchanges a pore-forming leucine residue located at the cytoplasmic entrance to the predicted EAAT/Glt_{Ph} anion conduction pathway and modifies EAAT2 anion channel open probabilities without significant alteration of transport rates. We interpret the effects on anion channel open probabilities by proposing that L46P affects the energy difference between intermediate transport conformation and open anion channels. The increased pore di-

ameter of L46P EAAT2 resulting in measurable gluconate[−] permeability suggests distinct positions of the transport domain in open anion channel conformation of WT and L46P EAAT2. Our findings provide additional experimental evidence for the recently proposed EAAT/Glt_{Ph} anion conduction mechanism (25).

Experimental Procedures

Heterologous expression of WT and mutant EAAT2

WT (kindly provided by Dr. M. Hediger, University of Bern, Switzerland) and mutant human EAAT2 were expressed as mYFP fusion protein as described previously (38). mYFP-EAAT2 fusion proteins have been intensively tested and been shown to be functionally indistinguishable from EAAT2 lacking the fluorescent protein (38). The L46P point mutation was introduced using PCR-based strategies, and the resulting pRcCMV-mYFP L46P EAAT2 was verified by restriction analysis and DNA sequencing. Two independent recombinants from the same transformation were examined and shown to exhibit indistinguishable functional properties. Transient transfection of HEK293T cells using the $Ca_3(PO_4)_2$ technique was performed as described previously (52, 53). Transfection rates were determined by manually counting transfected and untransfected cells on confocal images (2386 cells (three transfections) for WT and 2211 cells (three transfections) for L46P). Because macroscopic anion currents often exceeded 10 nA in cells transiently expressing L46P EAAT2, stable inducible cell lines expressing WT or mutant EAAT2 were generated by selection of Flp-In T-REx 293 cells (Invitrogen) transfected with pcDNA5/FRT/TO-EAAT2 and used either with or without induction with tetracycline (40, 52, 54).

Electrophysiology

Standard whole-cell or outside-out patch clamp recordings were performed using an HEKA EPC10 amplifier (HEKA Electronics, Lamprecht, Germany) as described (55). Borosilicate pipettes were pulled with resistances of 1.5–3.0 MΩ. To reduce voltage errors, we routinely compensated more than 80% of the series resistance by an analog procedure and excluded recordings with more than 10 nA maximum anion currents from the analysis. In experiments with outside-out patches, pipettes were covered with dental wax to reduce their capacitance. Currents were filtered at 2.9 kHz (−3 db) and digitized with a sampling rate of 50 kHz. Cells and patches were clamped to 0 mV for at least 4 s between sweeps.

To record anion currents (Figs. 2–4), the pipette solution contained (in mM) 115 Na/KNO₃, 2 MgCl₂, 5 EGTA, 10 HEPES, pH 7.4. The control extracellular bath solution contained (in mM) 140 NaNO₃ ± 1.0 L-glutamate. Under K^+ -bound homoexchange conditions, extracellular NaNO₃ was equimolar replaced by KNO₃. For reversal potential measurements (Fig. 4), several Na^+ -based external solutions supplemented with L-glutamate and varying NO₃[−]/gluconate[−] concentrations were used. We quantified contamination by background current by blocking EAAT2 currents by repeating the experiments with solutions lacking L-glutamate and supplemented with 0.1 mM DL-TBOA and corrected EAAT2 anion currents for the thus

obtained background currents. Relative gluconate permeabilities ($P_{\text{gluc}}/P_{\text{NO}_3}$) were calculated by fitting the Goldman-Hodgkin-Katz equation to concentration dependences of anion current reversal potentials (Fig. 4E):

$$E_{\text{rev}} = \frac{RT}{F} \ln \frac{P_{\text{gluc}}[\text{gluc}]_i + P_{\text{NO}_3}[\text{NO}_3]_i}{P_{\text{gluc}}[\text{gluc}]_o + P_{\text{NO}_3}[\text{NO}_3]_o} \quad (\text{Eq. 1})$$

For rapid application experiments, pipette solution contained (in mM) 115 NaSCN + 10 L-glutamate under glutamate-bound homoexchange conditions or 115 KSCN under transport conditions, and the bath solution 140 NaNO₃, 4 KCl, 2 CaCl₂, 1 MgCl₂, 5 HEPES, pH 7.4. To measure open probabilities under transport conditions or under homoexchange conditions, an internal solution, in which Na/KNO₃ were equimolarly substituted with K-gluconate, was used.

Kinetic modeling

To describe transient currents in symmetric K⁺, we used a kinetic model consisting of K⁺-bound and K⁺-free inward- and outward-facing conformations and an intermediate transport conformation, from which anion channel opening occurs (Fig. 6C). Model predictions of absolute open probabilities were fitted against experimentally determined values (Fig. 6). A genetic algorithm as implemented in the Python package DEAP was used for minimization of squared errors. Fits to WT and L46P data revealed exclusive alteration of anion channel opening/closing rates in L46P EAAT2.

Fast substrate application

For fast application of transport substrates, a piezo-driven system with a dual-channel theta glass (47, 56) tubing was used (Fig. 5) (MXPZT-300, Siskiyou, Grants Pass, OR, USA). At the end of each experiment, we removed the cell from the patch pipette and measured 20–80% rise times of our system by fast application of the used solution to the open pipette (56). We obtained values between 1.3 and 3.0 ms, with a mean value of 2.2 ± 0.4 ms (mean $\pm$ CI; $n = 30$).

Noise analysis

We used nonstationary noise analysis to determine single-channel current amplitudes of WT and L46P EAAT2 anion channels (Fig. 3) (41). Current variances were calculated from current differences in subsequent records during 300 subsequent voltage jumps (41) and subtracted the background noise measured at 0 mV. The 10% of the sweeps with the highest time-averaged variances were removed.

Ion channels usually generate a Lorentzian type of noise. Because open and closed states of an ion channel are binomially distributed, the amplitude and the time dependence of the current variance (σ^2) can be calculated by

$$\begin{aligned} \sigma(t)^2 &= N \times i^2 \times p(t) \times (1 - p(t)) + \sigma_{bg}^2 \\ &= I_{\text{mean}}(t) \times i - \frac{I_{\text{mean}}^2(t)}{N} + \sigma_{bg}^2 \end{aligned} \quad (\text{Eq. 2})$$

or after linear transformation

$$\frac{\sigma(t)^2 - \sigma_{bg}^2}{I_{\text{mean}}(t)} = i - \frac{I_{\text{mean}}(t)}{N} \quad (\text{Eq. 3})$$

with i being the unitary current amplitude, p the absolute open probability, N the total number of channels in the membrane, I_{mean} the mean macroscopic current amplitude, and σ_{bg}^2 the voltage-independent background noise. The y axis intercept from a linear regression of Equation 3 to the data points thus provides the unitary current amplitude, and the slope of the linear regression ($-N^{-1}$) the number of channels.

We normalized mean currents (I_{mean}) to the maximum current amplitudes (I_{max}) to account for differences in transporter expression and pooled normalized data from all cells expressing WT or mutant EAAT2 into single plots. We plotted ratios of current variances by non-normalized current amplitudes versus normalized current amplitudes for all examined cells (Fig. 3C). The regression error was assessed by bootstrap sampling simulations (43). For this, 50,000 bootstrap samples were resampled by randomly selecting data from the original datasets with replacement. We determined unitary current amplitudes for all bootstrap samples and the distribution of these values was plotted in a histogram for visual inspection (Fig. 3D). The reported errors of the single-channel currents are the confidence intervals of this parameter of the regression of all bootstrap samples.

Confocal imaging

Images were acquired 24–36 h after transfection with a Leica TCS SP5 II inverted microscope (Mannheim, Germany) using a 63 $\times$ oil immersion objective from living cells in PBS containing Ca²⁺ and Mg²⁺ at room temperature (22–24°C) (57). mYFP (monomeric yellow fluorescence proteins) fluorophores were excited with a 488-nm argon laser. Confocal images were assembled for publications in ImageJ (58).

Biochemical analysis

Membrane proteins were solubilized with 0.4% DDM after membrane isolation from transfected HEK293T using hypotonic and hypertonic solutions (59). Total protein concentrations were determined using the BCA assay, and cleared lysates were then denatured for 15 min at room temperature in SDS sample buffer and electrophoresed in parallel with fluorescent mass markers (Dual Color, Bio-Rad) on 10% SDS-polyacrylamide gradient gels. YFP-tagged proteins were visualized by scanning the wet PAGE gels with a fluorescence scanner (Typhoon, GE Healthcare, München, Germany). Individual bands were quantified with ImageJ (53).

Data analysis

Data analysis was performed using a combination of FitMaster (HEKA), self-written programs in Python (Continuum Analytics), Origin (OriginLab), SigmaPlot (Systat Software), and

Excel (Microsoft) software. All data are presented as mean $\pm$ 95% confidence interval or S.D. as indicated. For fast application and noise analysis, S.D. was assessed by bootstrap sampling simulations with 50,000 bootstrap samples, randomly selected from the original dataset. In boxplot diagrams, *boxes* represent 25.75 percentiles, *whiskers* indicate 95% confidence intervals, *straight lines* show the median value, and *dotted lines* show the mean value. Mean values were compared using two-way analysis of variance with Holm-Sidak post hoc test. Data are significant different with $p \leq 0.05$ (*), $p \leq 0.01$ (**), or $p \leq 0.001$ (***).

Data availability

All relevant data and scripts are available at <https://github.com/dkortzak/EAAT2-mutation-affecting-anion-currents>.

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Abbreviations—The abbreviations used are: EAAT, excitatory amino acid transporter; CI, confidence interval.

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