

Pyridoxal kinase inhibition by artemisinins down-regulates inhibitory neurotransmission

Vikram Babu Kasaragod^{a,1,2,3}, Anabel Pacios-Michelena^{a,1}, Natascha Schaefer^b, Fang Zheng^c, Nicole Bader^a, Christian Alzheimer^c, Carmen Villmann^b, and Hermann Schindelin^{a,3}

^aInstitute of Structural Biology, Rudolf Virchow Center for Integrative and Translational Bioimaging, University of Würzburg, 97080 Würzburg, Germany; ^bInstitute for Clinical Neurobiology, University of Würzburg, 97078 Würzburg, Germany; and ^cInstitute of Physiology and Pathophysiology, Friedrich Alexander University Erlangen-Nürnberg, 91054 Erlangen, Germany

Edited by Lily Yeh Jan, University of California, San Francisco, CA, and approved October 26, 2020 (received for review May 3, 2020)

The antimalarial artemisinins have also been implicated in the regulation of various cellular pathways including immunomodulation of cancers and regulation of pancreatic cell signaling in mammals. Despite their widespread application, the cellular specificities and molecular mechanisms of target recognition by artemisinins remain poorly characterized. We recently demonstrated how these drugs modulate inhibitory postsynaptic signaling by direct binding to the postsynaptic scaffolding protein gephyrin. Here, we report the crystal structure of the central metabolic enzyme pyridoxal kinase (PDXK), which catalyzes the production of the active form of vitamin B6 (also known as pyridoxal 5'-phosphate [PLP]), in complex with artesunate at 2.4-Å resolution. Partially overlapping binding of artemisinins with the substrate pyridoxal inhibits PLP biosynthesis as demonstrated by kinetic measurements. Electrophysiological recordings from hippocampal slices and activity measurements of glutamic acid decarboxylase (GAD), a PLP-dependent enzyme synthesizing the neurotransmitter γ -aminobutyric acid (GABA), define how artemisinins also interfere presynaptically with GABAergic signaling. Our data provide a comprehensive picture of artemisinin-induced effects on inhibitory signaling in the brain.

artemisinins | pyridoxal kinase | pyridoxal phosphate (PLP) | GABA biosynthesis | inhibitory neurotransmission

Pyridoxal 5'-phosphate (PLP) is the active form of vitamin B6. In humans, PLP biosynthesis is catalyzed by pyridoxal kinase (PDXK), a member of the ribokinase superfamily. PDXK utilizes inactive forms of vitamin B6 (pyridoxal [PL], pyridoxine, and pyridoxamine) and ATP as substrates, producing PLP along with the byproduct ADP. The corresponding reaction proceeds via a random substrate addition reaction mechanism (1) in which PLP biosynthesis takes place by transferring the γ -phosphate of ATP to the 5'-OH group of the B6 vitamers, in a process assisted by divalent metal ions such as Zn^{2+} and Mg^{2+} (2) (Fig. 1A). PLP serves as the essential active site component for more than 160 distinct human enzymatic activities (3) catalyzing crucial cellular processes such as detoxification reactions and multiple metabolic processes including amino acid, carbohydrate, and lipid metabolism. PLP-dependent enzymes also participate in neurotransmitter biosynthesis including the inhibitory neurotransmitters γ -aminobutyric acid (GABA) and glycine (4–6), which are synthesized by glutamic acid decarboxylase (GAD) and serine hydroxymethyl transferase (SHMT), respectively. Vitamin B6 deficiency has been implicated in multiple neurological, psychiatric, and internal disorders possibly including even diabetes, cancer, and autism (3), thus underpinning the importance of a finely tuned PLP biosynthesis.

Recently, PDXK was identified as one of the mammalian targets of the antimalarial drug artemisinin (7). Artemisinincontaining plant extracts have been used in traditional Chinese medicine for the treatment of malaria (8). Chemically, these small molecules are sesquiterpene lactones with an unusual endoperoxide bridge. Artemisinin and its semisynthetic derivatives artemether and artesunate (collectively referred to as artemisinins), in combination with quinones such as mefloquine and lumefantrine, nowadays represent the standard drug combinations used to treat malaria caused by *Plasmodium falciparum* (9). In addition to their antiprotozoan activities, these drugs have also been pharmacologically observed to regulate the activities of a variety of mammalian cellular processes, some of which are deregulated in various types of cancer (10, 11). Recently, it was discovered that artemisinins also modulate the differentiation of pancreatic T α cells by inducing the transdifferentiation of glucagon-producing T α cells into insulin-secreting T β cells, thus suggesting an antidiabetic activity of artemisinins (7). However, two subsequent studies contradicted this observation, thus questioning the potential clinical application of these compounds in the treatment of diabetes (12, 13).

Until recently, in the absence of a single protein crystal structure in complex with artemisinins (neither a plasmodial nor a mammalian protein), the detailed framework describing the target recognition by these small molecules remained enigmatic. The first molecular insights into artemisinin recognition by a target protein were derived by us from crystal structures of the C-terminal domain of the moonlighting protein gephyrin (GephE) in complex with two artemisinin derivatives, artesunate and artemether (14). Gephyrin is

Significance

Information processing in the central nervous system relies on chemical synapses where neurotransmitters such as the inhibitory neurotransmitter γ -aminobutyric acid (GABA) are released at presynaptic nerve endings. GABA is synthesized by glutamic acid decarboxylase (GAD), an enzyme requiring pyridoxal 5'phosphate (PLP or vitamin B6) as cofactor, the latter being synthesized by pyridoxal kinase (PDXK). Here, we show that the antimalarial drug artemisinin inhibits PDXK and describe the structural basis of this inhibition. The decrease in PLP production reduces the amount of GABA being produced, which, in turn, impacts the efficacy of GABAergic transmission. This study combined with our previous data sheds light on how artemisinins can influence inhibitory synaptic transmissions both presynaptically, as described here, and postsynaptically.

The authors declare no competing interest.

This article is a PNAS Direct Submission.

Published under the PNAS license.

¹V.B.K. and A.P.M. contributed equally to this work.

This article contains supporting information online at https://www.pnas.org/lookup/suppl/ doi:10.1073/pnas.2008695117/-/DCSupplemental.

First published December 14, 2020.

Author contributions: V.B.K., A.P.M., N.S., F.Z., C.A., C.V., and H.S. designed research; V.B.K., A.P.M., N.S., F.Z., N.B., and C.V. performed research; V.B.K., A.P.M., N.S., F.Z., N.B., C.V., and H.S. analyzed data; and V.B.K., C.A., C.V., and H.S. wrote the paper.

²Present address: Neurobiology Division, Medical Research Council Laboratory of Molecular Biology, Cambridge CB2 0QH, United Kingdom.

³To whom correspondence may be addressed. Email: vkasaragod@mrc-lmb.cam.ac.uk or hermann.schindelin@virchow.uni-wuerzburg.de.

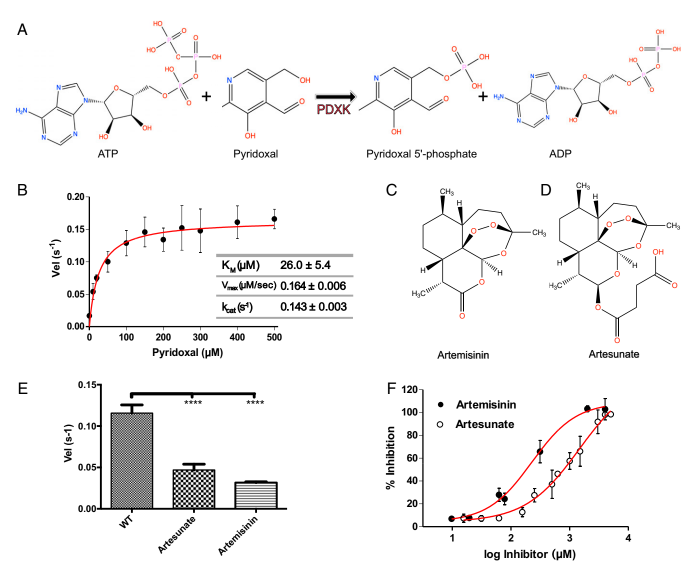


Fig. 1. Biochemical basis of PDXK inhibition by artemisinins. (*A*) Schematic representation of the reaction catalyzed by PDXK. (*B*) Michaelis–Menten curve derived for the enzymatic activity of recombinantly purified PDXK. (*C* and *D*) Chemical structures of artemisinin (*C*) and artesunate (*D*). (*E*) Enzymatic activity of wild-type PDXK (WT-PDXK) in the absence and presence of artemisinin derivatives at a concentration of 1.5 mM (artesunate) and 156 μ M (artemisinin), respectively. Data are presented as mean \pm SEM (*P* values are **P* < 0.05; ***P* < 0.01; ****P* < 0.001; *****P* < 0.0001) (one-way ANOVA test). (*F*) Inhibition curves of PDXK by artemisinin and artesunate used to derive the corresponding IC₅₀ values.

the principal scaffolding protein at inhibitory postsynaptic specializations and also catalyzes the final two steps of the evolutionarily conserved molybdenum cofactor (Moco) biosynthesis (15–17). Structures of the GephE–artemisinin complexes demonstrated that artemisinins specifically target the universal receptor binding pocket of this moonlighting protein, without altering its enzymatic activity, thus inhibiting critical interactions of gephyrin with GABA type A receptors (GABA_ARs) and glycine receptors (GlyRs). As an important functional consequence, artemisinins modulate inhibitory neurotransmission in a gephyrin-dependent manner. In addition to gephyrin, various proteins were identified as putative targets of artemisinins in pancreatic cells, including the central metabolic enzyme PDXK (7), yet the molecular mechanisms underlying the modulation of these targets by artemisinins remained unknown.

Here, we determined the 2.4-Å resolution crystal structure of mouse pyridoxal kinase (mPDXK) in complex with artesunate, a succinate derivative of artemisinin. The artesunate binding site partially overlaps with the substrate (PL)/product (PLP) binding site, thus suggesting a drug-induced inhibitory effect. Enzymatic activity assays in vitro indeed revealed a significant inhibition of

e (PL)/product (PLP) binding d inhibitory effect. Enzymatic

PLP production in the presence of artemisinins with K_i values in the high micromolar range. Electrophysiological recordings and measurements of GABA biosynthesis suggest that artemisinins exert their effect by down-regulating the activity of PLP-dependent enzymes such as GAD. Taken together, our data define the molecular basis for the inhibition of PDXK by artemisinins and their consequences at the presynaptic terminals of inhibitory postsynapses and extend our current understanding of the artemisinin-induced modulation of inhibitory neurotransmission beyond gephyrin.

Results

Artemisinins Inhibit PDXK. To derive the oligometric state of recombinantly purified mPDXK, we performed multiangle laser light scattering coupled to size exclusion chromatography (SEC-MALLS). The experiments showed that the protein is a dimer in solution (*SI Appendix*, Fig. S1), as has been reported for the human and also prokaryotic PDXK homologs (18). Next, we measured the enzymatic activity of mPDXK by directly monitoring PLP production in a photometric assay revealing a K_M of $26.0 \pm 5.4 \mu$ M, a V_{max} of $0.1640 \pm 0.006 \mu$ M/s, and a k_{cat} of $0.1436 \pm 0.003 \text{ s}^{-1}$ for the

substrate PL in the presence of 1 mM of ATP (Fig. 1*B*), which is in line with reported K_M values (3 to 50 μ M) for PDXK (19–25).

To understand the effect of artemisinins on the enzyme, we performed the activity assays in the presence of two artemisinins, the parental compound artemisinin and the succinate derivative artesunate (Fig. 1 C and D). The determination of the turnover rates (velocity [Vel]) displayed a highly significant inhibition in mPDXK activity in the presence of artemisinins with observed reductions to 0.032 ± 0.001 and 0.047 ± 0.007 s⁻¹ for artemisinin and artesunate, respectively. Compared to the turnover rate of the enzyme in the absence of these drugs (0.116 \pm 0.01 s⁻¹) (Fig. 1E) this corresponds to an approximately threefold decrease. The enzymatic or turnover velocity, Vel, is defined here as the mean number of product molecules generated by a single enzyme per unit time. Statistical analyses revealed a significant reduction in enzymatic activity in the presence of the artemisinins and via a Dixon plot analysis K_i values of 120 ± 2.4 and $1250 \pm 4.7 \ \mu M$ were derived for artemisinin and artesunate, respectively (SI Appendix, Fig. S2). To further characterize the inhibitory properties of artemisinins we also determined the IC_{50} for both compounds. While artesunate displayed an IC₅₀ value of 1,445 \pm 1.4 μ M, artemisinin was approximately sixfold more potent with an IC₅₀ of 229 \pm 1.3 μ M (Fig. 1F). These results were in contrast to what we observed earlier for gephyrin, the postsynaptic target of the artemisinins, where the highest affinity and the higher stabilization of the C-terminal E domain were observed with artesunate.

Owing to a superior solubility of artesunate (approaching ~ 10 mM at moderate dimethyl sulfoxide (DMSO) concentrations), all subsequent in vitro experiments with PDXK including our crystallization trials were carried out with artesunate.

Structural Basis for the Inhibition of PDXK by Artemisinins. To gain insights into the mechanism of inhibition at the atomic level we determined three crystal structures of mPDXK. First, we derived the crystal structure of mPDXK in its apo-state and in complex with ATP γ S, one of the substrates of the enzyme. These structures were solved by molecular replacement (MR) with the structure of human apo-PDXK as the search model. The apo and the mPDXK-ATP γ S structures were refined in space group C2, containing two dimers in the asymmetric unit, to resolutions of 2.45 and 2.9 Å, respectively (Table 1 and *SI Appendix*, Fig. S3 *A* and *B*). The overall architecture of mouse apo-PDXK shares high structural similarity with its human ortholog (26, 27) as reflected in root-mean-square (rms) deviations of 0.84 Å (Protein Data Bank [PDB]: 2YXT; human apo-PDXK) after superposition of all C α atoms.

Closer inspection of the nucleotide binding pocket revealed that ATP γ S binding is directly mediated by Val226, which forms a hydrogen bond with the adenine of the nucleotide through its main chain carbonyl oxygen and residues Thr186 and Thr233 as well as Asp118 and Asn150, which coordinate the ATP analog through interactions with the α - and β -phosphates of its triphosphate moiety, respectively (*SI Appendix*, Fig. S3*C*). There were no significant conformational changes in the binary complex compared to the mouse apo-structure as reflected in an rms deviation of 0.45 Å for all C α atoms with minimal structural rearrangements in the ATP binding pocket (*SI Appendix*, Fig. S3*D*). A comparison of PDXK sequences derived from organisms representing different evolutionary levels revealed that all residues, which are crucial for the binding of the nucleotide, are strictly conserved (*SI Appendix*, Fig. S4).

To gain insights into the mechanism of artemisinin inhibition we determined the crystal structure of mPDXK in complex with ATP γ S and artesunate (Fig. 2 and Table 1). This structure was obtained by soaking artesunate into preexisting binary mPDXK-ATP γ S crystals. After MR with the apo-structure, in addition to the clear density for ATP γ S (Fig. 2*B*), strong difference density in close proximity to the substrate-binding pocket was also observed (Fig. 2*D*), which allowed us to unambiguously model the bound artesunate. Surprisingly, this density was observed in only one of the four molecules present in the asymmetric unit. The absence of artesunate in the other monomers may be due to the involvement of these protomers in crystal contacts, thus preventing artesunate binding when soaking the compounds into preexisting crystals.

The fact that all three structures reported here belong to the same space group with similar unit cell parameters and essentially identical crystal packing allowed for a meaningful comparative analysis. The overall architecture of the mPDXK-ATPγS-artesunate structure is identical to that of the apo and binary mPDXK-ATPyS structures; a superposition of the Ca atoms of these two complexes revealed rms deviations of 0.51 and 0.30 Å for the apo and ATP γ S-bound structures, respectively. Binding of the substrate analog ATP γ S was mediated by the same residues described for the binary mPDXK-ATPyS complex (Fig. 2C and SI Appendix, Fig. S5). A closer inspection of the artesunate-binding pocket revealed that drug binding is mainly mediated by Val41, Thr47, and also Trp52, which generate a hydrophobic pocket that binds artesunate with a buried surface area of 364 Å^2 compared to a total surface area of the drug of 538 Å^2 . In particular, artesunate is sandwiched in between two aromatic residues, Phe43 and Tyr84, which stabilize artesunate through van der Waals interactions. In addition to the hydrophobic interactions, the carboxylate moiety of artesunate comes into proximity of the guanidinium group in the side chain of Arg86, which potentially stabilizes the interactions through electrostatic contacts. Finally, Asp87 favors artesunate binding through a hydrogen bond (2.5 Å) between its side chain and the carboxylate of artesunate, assuming one of these carboxylates is protonated (Fig. 2E).

An analysis of the mPDXK-ATPyS-artesunate structure showed that the ATP binding pocket is in relatively close proximity to the bound artesunate at a distance of ~21 Å, as measured between the $C\alpha$ atoms of Phe237 in the ATP binding pocket and Phe43 in the artesunate binding pocket. A comparison of the ternary mPDXK-ATPyS-artesunate and binary mPDXK-ATPyS structures suggested that binding of artesunate neither induced any significant rearrangements in the conformation of the nucleotide nor in the residues mediating its binding, in line with the independent binding of the two ligands (SI Appendix, Figs. S5 and S6). Interestingly, the binding pocket of artesunate, like that of ATP mentioned earlier, is highly conserved (SI Appendix, Fig. S7). To get additional information regarding the mode of inhibition, we also compared our ternary structure with PDXK-PLP structures. Strikingly, when we superimposed the already reported ternary HsPDXK-ATP-PLP (PDB: 3KEU) structure with our ternary complex, a critical partial overlap between the tricyclic ring system of artesunate and the pyridine ring of the product PLP was uncovered (Fig. 3Aand B), which would result in severe van der Waals repulsions for the C2 and C3 atoms of PLP and the C2 and C3 atoms of artesunate, if bound simultaneously. This clearly constitutes a major reason for the inhibition of the PDXK. Moreover, an analysis of the surface properties of the ternary structure revealed a tunnel, which is leading from the protein surface to the distal end of the ATP binding pocket spanning a length of \sim 38 Å, which is blocked near its entrance by artesunate. A blockade of this tunnel, in turn, may prevent an efficient turnover of the enzyme. Taken together, these data illustrate the structural basis for the inhibition of mPDXK by artemisinins (Fig. 3 *C*–*E*).

Mapping of the Artemisinin Binding Pocket. To validate the observations derived from the crystal structures, we performed sitedirected mutagenesis experiments of residues located in the

Table 1. Data collection and refinement statistics

Data collection	PDXK-apo	ΡΟΧΚ-ΑΤΡγS	PDXK-ATPγS–artesunate
Space group		C2	
a, b, c (Å)	279.13, 53.43, 109.37	278.60, 53.02, 109.85	279.38, 53.04, 110.15
α, β, γ (°)	90, 90.00, 90	90, 91.75, 90	90, 91.64, 90
Resolution (Å)	47.32 to 2.45 (2.53 to 2.45)	47.16 to 2.9 (3.03 to 2.9)	47.20 to 2.4 (2.46 to 2.4)
<i>R</i> sym*	0.098 (0.75)	0.10 (0.739)	0.084 (1.069)
Rpim [†]	0.093 (0.70)	0.068 (0.488)	0.053 (0.704)
CC _{1/2}	0.993 (0.572)	0.995 (0.600)	0.996 (0.541)
< <i>l</i> / σ <i>l</i> > [‡]	9.1 (1.6)	9.3 (1.6)	8.1 (1.1)
Completeness (%)	95.6 (99.7)	99.7 (99.7)	99.7 (99.6)
Redundancy	3.2 (3.5)	3.4 (3.2)	3.4 (3.1)
Reflections used in refinement	57,174 (5,965)	36,057 (3,532)	63,558 (6,235)
Refinement			
R-work [¶]	0.2162 (0.3042)	0.2341 (0.3378)	0.2088 (0.3271)
R-free [§]	0.2596 (0.3237)	0.2548 (0.3366)	0.2535 (0.3851)
No. of nonhydrogen atoms	9,715	9,862	9,890
Macromolecules	9,411	9,606	9,445
Ligands	116	232	304
Solvent	188	24	141
RMS (bonds)	0.002	0.005	0.002
RMS (angles)	0.53	0.73	0.54
Ramachandran favored (%) [#]	96.83	96.99	96.76
Ramachandran allowed (%)	3.17	3.01	3.24
Ramachandran outliers (%)	0.00	0.00	0.00
Average B-factor (Å ²)	57.79	95.87	69.33
Protein	57.62	95.89	68.94
Ligands	74.74	95.54	84.27
Solvent	55.93	91.85	63.08

Although the β angle of the apo-structure is 90.00, the crystals display monoclinic and not orthorhombic symmetry. Numbers in parentheses refer to the respective highest-resolution data shell in each dataset.

* $R_{sym} = \sum_{hkl} \sum_i |I_i - \langle I \rangle |\Sigma_{hkl} \sum_i I_i$, where I_i is the *i*th measurement and $\langle I \rangle$ is the weighted mean of all measurements of *I*.

 ${}^{\dagger}R_{\text{pim}} = \sum_{hkl} 1/(N-1)^{1/2} \sum_{i} I_{i}(hkl) - I_{i}(hkl) |\Sigma_{hkl} \sum_{i} (hkl), \text{ where } N \text{ is the redundancy of the data and } I_{i}(hkl) \text{ the average intensity.}$

 $^{*}</l/\sigma/>$ indicates the average of the intensity divided by its SD.

 $||\mathbf{R}_{work} = \sum_{hkl} ||\mathbf{F}_o| - |\mathbf{F}_c|| / \sum_{hkl} ||\mathbf{F}_o|$, where \mathbf{F}_o and \mathbf{F}_c are the observed and calculated structure factor amplitudes.

 ${}^{\$}R_{free}$ same as R for 5% of the data randomly omitted from the refinement. The number of reflections includes the R_{free} subset.

[#]Ramachandran statistics were calculated with MolProbity.

artesunate binding pocket (Fig. 3F) and tested these mutants for PLP production in the presence and absence of artesunate (Fig. 3 G and H). First, we analyzed the mutants through SEC-MALLS, which revealed that all variants retained their dimeric state in solution as observed for the wild-type (WT) protein (SI Appendix, Fig. S8). Among the residues being investigated, we mutated Val41 and Phe43, which are involved in mediating the binding of artesunate through hydrophobic interactions to either introduce steric interference or alter the polar properties of the binding pocket, respectively. The V41W and F43R mutants significantly lowered the turnover rates $(0.04 \pm 0.006 \text{ s}^{-1} \text{ for V41W})$ and $0.016 \pm 0.001 \text{ s}^{-1}$ for F43R), even in the absence of artesunate in comparison to the WT ($0.080 \pm 0.004 \text{ s}^{-1}$). This can be easily explained by the fact that these residues also mediate binding of PL and thus play a role in the regular enzymatic turnover of the protein. In contrast, mutation of Arg86, the residue involved in the long-range electrostatic interaction with the carboxylate of artesunate, to the bulky aromatic side chain of Trp did not alter the activity of mPDXK in the absence of artesunate, as demonstrated by its turnover rate of 0.08 \pm 0.005 s^{-1} , which is virtually identical to that of the WT (Fig. 3G).

Next, we further analyzed the catalytic activity of the mutants in the presence of artesunate (Fig. 3*H* and *SI Appendix*, Fig. S9). The V41W and F43R variants did not result in significant changes in the turnover rates of the enzyme in the presence of artesunate $(0.044 \pm 0.009 \text{ s}^{-1} \text{ for V41W} \text{ and } 0.014 \pm 0.005 \text{ s}^{-1} \text{ for}$ F43R compared to 0.040 ± 0.006 and $0.016 \pm 0.001 \text{ s}^{-1}$, respectively, in its absence), which is in line with artesunate binding being abolished in both variants (SI Appendix, Fig. S9 C and D). As expected, a similar trend was observed in the case of the V41W/ F43R double mutant with turnover rates of 0.024 ± 0.002 s⁻¹ in the presence of artesunate compared to $0.019 \pm 0.004 \text{ s}^{-1}$ in its absence (SI Appendix, Fig. S9E). An identical behavior was observed in the case of GephE where mutation of a crucial aromatic residue (Phe330) to Ala completely abolished artemisinin binding (14). In contrast, when we compared the activity of the R86W variant in the absence $(0.080 \pm 0.005 \text{ s}^{-1})$ and presence $(0.038 \pm 0.008 \text{ s}^{-1})$ of the drug, a significant reduction in enzymatic activity was observed (SI Appendix, Fig. S9B). Thus, although R86 is involved in an electrostatic interaction as revealed by the crystal structure, the mutational analysis demonstrated that the inhibition potency of artesunate is retained even in the absence of this interaction. This observation is in contrast to the GephE structure where the replacement of Arg (Arg653 in gephyrin) with the bulkier aromatic Trp prevented artesunate binding (14). Thus, our structures help to define the molecular signatures of artemisinin binding pockets, which may aid in the future identification of target sites, especially by in silico approaches.

Artemisinins Inhibit GABA Biosynthesis and Down-Regulate GABAergic Neurotransmission. To understand whether the functional consequences of our biochemical and structural analyses correlate with a physiological scenario, whole-cell voltage-clamp recordings (Fig. 4) from CA1 pyramidal cells in hippocampal slices were performed. We determined the properties of GABAergic miniature inhibitory postsynaptic currents (mIPSCs) in the absence and

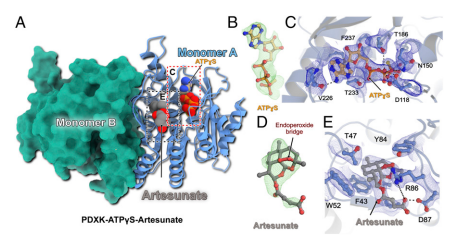


Fig. 2. Structure of the ternary PDXK-ATP γ S-artesunate complex. (A) Overall architecture of the ternary complex. One monomer is shown in cartoon representation with the bound ligands in CPK representation, while the second monomer is shown as a surface in green. (B) F_o - F_c omit electron density for the bound ATP γ S contoured at an rms deviation of 3. (C) Enlarged view of the ATP γ S binding pocket. The bound ligand and residues, which are crucial for ligand binding, are shown in ball-and-stick representation. The SIGMAA-weighted $2F_o$ - F_c electron density for the bound ligand and surrounding residues is contoured at an rms doi 1. Critical protein–ligand interactions are highlighted. (D) F_o - F_c omit electron density for the bound artesunate contoured at an rms deviation of 3. (E) Enlarged view of the artesunate-binding pocket. The bound ligand and residues, which are crucial for ligand binding, are shown in ball-and-stick representation. SIGMAA-weighted $2F_o$ - F_c electron density for the bound artesunate contoured at an rms deviation of 3. (E) Enlarged view of the artesunate-binding pocket. The bound ligand and residues, which are crucial for ligand binding, are shown in ball-and-stick representation. SIGMAA-weighted $2F_o$ - F_c electron density for artesunate and interacting residues is contoured at an rms of 1. Critical protein–ligand interactions are highlighted. The bound ligand and residues, which are crucial for ligand binding, are shown in ball-and-stick representation. SIGMAA-weighted $2F_o$ - F_c electron density for artesunate and interacting residues is contoured at an rms of 1. Critical protein–ligand interactions are highlighted.

presence of artemisinin (10 and 30 µM, Fig. 4 B and C) as well as artesunate at a concentration of 50 µM (Fig. 4A), which was shown to deregulate the postsynaptic GABAAR-gephyrin clustering. Artesunate treatment induced a significant reduction in the mIPSC amplitudes from 45.4 ± 4.3 to 35.7 ± 3.9 pA (P = 0.0001), while the frequency decreased from 5.9 ± 0.7 to 4.6 ± 0.7 Hz (P = 0.0008, n = 8 from four mice) at the tested concentration (Fig. 4A, D, and E). In contrast, artemisinin down-regulated mIPSC amplitudes already at 10 μ M (from 52.7 \pm 1.9 to 38.8 \pm 2.6 pA, n = 7from four mice, P = 0.001, paired t test; Fig. 4 B and D). We attribute the changes in the amplitude to the artemisinin-induced disruption of postsynaptic GABAAR-gephyrin complexes. While a significant reduction in amplitudes was retained at a higher concentration (30 μ M), we also observed a concomitant decrease in mIPSC frequency from 5.1 ± 0.6 to 3.9 ± 0.5 Hz (n = 7 from five mice, P = 0.012; Fig. 4 C and E) in the presence of artemisinin. In addition, artemisinin and/or artesunate altered mIPSC kinetics with slower rise and decay times (Fig. 4 F and G). In both cases, a significant decrease in mIPSC frequency is particularly noteworthy in the context of this study as it reflects changes in the presynaptic terminals; e.g., it would be in line with a reduced synthesis of the neurotransmitter GABA (28).

To evaluate whether the changes in the mIPSC frequencies observed were due to changes in the cellular concentration of PDXK, we first analyzed its expression levels in hippocampal neurons at day in vitro (DIV) 14 (Fig. 5 A and B), as PDXK activity is required for the activity of GAD by providing its cofactor PLP. These studies revealed that the PDXK expression level was unaffected by artemisinin treatment. Next, we investigated whether the observed mIPSC frequencies were due to changes in GABA availability upon alterations in the activity of GAD, the GABA synthesizing enzyme. Following treatment with artemisinin at different concentrations, no changes in the overall GAD65 expression level were observed (Fig. 5 C and D). In contrast to GAD67 which is constitutively active, GAD65 is transiently active when there is an increased demand of GABA. Moreover, it was shown that GAD65 is activated by phosphorylation (29, 30). To investigate whether treatment with artemisinin in vitro has an impact on the amount of active phosphorylated GAD65, immunoprecipitation studies were performed. Using hippocampal neurons at DIV14 in culture treated with increasing artemisinin concentrations (1, 3, 10, and 30 µM), a reduction in the levels of phosphorylated GAD65 protein was detected compared to the DMSO-treated control samples (1 μ M 55 \pm 9% compared to DMSO 100%, n = 4, P = 0.108; 3 μ M 48 \pm 7%, $n = 4, P = 0.059; 10 \ \mu\text{M} 55 \pm 11\%, n = 4, P = 0.11; 30 \ \mu\text{M} 30 \pm 10\%$ 6%, n = 2, P = 0.09); however, the data fell just short of reaching statistical significance (Fig. 5 C and E). As a qualitative measure, we also stained hippocampal neurons for PDXK, GAD, and the postsynaptic marker gephyrin, which did not reveal any noticeable differences in the expression of these marker proteins (SI Appendix, Figs. S10 and S11). Finally, to check whether the frequency changes observed in the electrophysiology measurements and the down-regulation observed in the active form of GAD correlate with GABA biosynthesis, we measured the amount of this neurotransmitter in primary hippocampal neurons (DIV14). GABA levels were quantified by the classical ninhydrin reaction (SI Appendix, Fig. S12) by measuring the fluorescence emission of the resulting adduct at 450 nm and calibrating it with a GABA standard. Remarkably, this analysis revealed a significant reduction in the amount of GABA production (5.4 \pm 0.8, 3.2 \pm 0.9, and 3.6 \pm 0.98 µg GABA per milligram protein per hour) in hippocampal neurons treated with artemisinin at concentrations of 3, 10, and 30 µM, respectively (P = 0.0054, 0.0007, and 0.0002 against DMSO measurements)and P = 0.018, 0.0087, and 0.0047 against untreated hippocampal measurements for 3, 10, and 30 µM artemisinin concentrations) (Fig. 5F). In comparison, untreated samples resulted in levels of $9.8 \pm 1.5 \ \mu g$ GABA per milligram protein per hour. The observed perturbation on the presynaptic side is therefore primarily due to the direct effect of artemisinins on the biosynthesis of PLP, which results in a reduced production of this cofactor, which is required by the GAD enzyme to produce the neurotransmitter GABA (Fig. 5G), while a slight augmentation of this decrease may be due to a reduction in the levels of the phosphorylated form of GAD65.

Discussion

Despite their widespread clinical application as antimalarial drugs, and despite their known effects on various cellular pathways in mammals, the molecular mechanisms of how artemisinins affect cellular pathways are still only poorly understood.

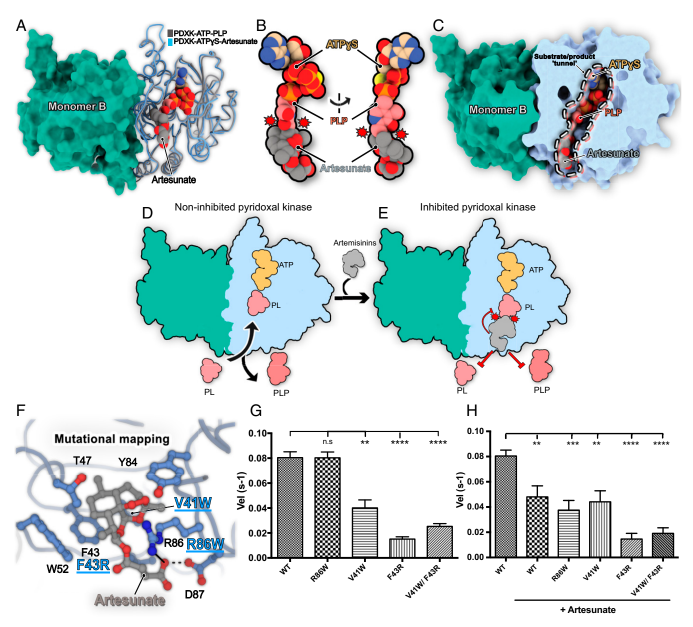


Fig. 3. Structural basis for PDXK inhibition by artemisinins. (*A*) Superposition of the crystal structures of the ternary murine PDXK-ATP₇S-artesunate (this study) and the human PDXK-ATP-PLP (PDB: 3KEU) complexes. Carbon atoms of artesunate are shown in gray, those of PLP in pink, and those of ATP₇S in beige; the other atoms are colored in red (oxygen), blue (nitrogen), orange (phosphorous), and yellow (sulfur). Only ATP₇S of the artesunate complex is shown to reduce visual complexity. (*B*) Enlarged view of the ligand binding pockets of ATP, artesunate, and PLP displaying the partial overlap (indicated by red circles with spikes) between PLP and artesunate. (*C*) Cutaway view of the superimposed PDXK-ATP₇S-artesunate and PDXK-ATP₇S-PLP structures displaying how artesunate binding blocks the substrate tunnel, which may also impair enzyme turnover. (*D* and *E*) Schematic representation of the structural basis for inhibition of PDXK activity by artemisinins. (*F*) Mutational mapping of the artesunate binding pocket with the investigated mutants highlighted in blue. (*G* and *H*) Comparison of the turnover rates of PDXK variants and the WT in the absence (*G*) and presence (*H*) of artesunate (**P* < 0.05; ***P* < 0.01; *****P* < 0.001) (paired *t* test).

Artemisinins can efficiently cross the blood-brain barrier (31) and, strikingly, administrations of high levels of artemisinins are accompanied by severe neurotoxic side effects (32–34). Recently, we were able to derive the first protein-artemisinin structure by X-ray crystallography at 1.5 Å resolution, namely that of the inhibitory postsynaptic scaffolding protein gephyrin in complex with artesunate and artemether (14). Here, we successfully validated and elucidated the mechanism underlying yet another mammalian artemisinin target, the critically important metabolic enzyme PDXK.

Our structural studies demonstrate a competition between the substrate pyridoxal and artemisinins, in line with the observed

inhibition of the enzyme derived from kinetic data. As artesunate targets the same binding pocket identified previously for the interaction of (R)-roscovitine with PDXK (35, 36) and for the neurotoxins ginkgotoxin and theophylline (37), our structure suggests that the neurotoxicity induced by artemisinins could be due, at least in part, to their binding to PDXK and the resulting inhibition of its activity.

The presynaptic effect of artemisinin and, at higher concentrations, also for artesunate in our electrophysiological recordings correlates nicely with the down-regulation of PDXK activity and can be extended toward glycine, the other major inhibitory neurotransmitter. This neurotransmitter is synthesized by SHMT,

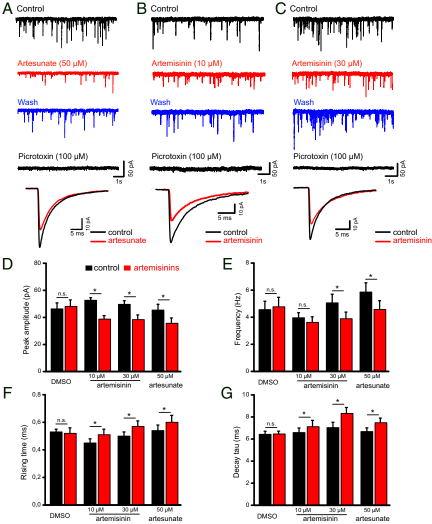


Fig. 4. Impact of artemisinins on electrophysiological recordings of hippocampal slices. (A–C) Representative voltage-clamp recordings of GABA_A receptormediated mIPSCs from mouse CA1 pyramidal cells in hippocampal slices were collected before (control), during (10 to 15 min), and after (wash) artesunate (50 μ M, A) or artemisinin (10 μ M, B and 30 μ M, C) application. Picrotoxin was applied to verify the GABAergic origin of these events. Superimposed traces (A–C, *Bottom*) are averaged events before and during artemisinin treatment. (D–G) Quantifications of how artemisinins affect mIPSC amplitudes (D), frequencies

again in a strictly PLP-dependent fashion. Thus, we predict a similar electrophysiological behavior with decreased frequencies at glycinergic synapses as observed for GAD and GABA levels. With respect to the postsynaptic counterparts, we have already demonstrated a decrease in glycinergic currents following artemisinin treatment (14). Structurally, GlyRs, via their β subunit, also depend on the identical binding pocket in GephE for their postsynaptic clustering as do GABA_ARs. Our GephE–artesunate/artemether structures demonstrated that artemisinins block the N-terminal three-residue stretch from the core binding motifs of the GlyR and GABA_ARs to mediate the inhibition (³⁹⁸FSI⁴⁰⁰ of GlyR β subunit vs. ³⁹⁸FNI⁴⁰⁰ of GABA_AR α 3 subunit). Therefore, we can extrapolate these results and predict a similar down-regulation of receptor clustering at glycinergic postsynapses.

(E), and kinetics (F and G). *P < 0.05, paired t-test.

Thus, the data presented here extend our current understanding of how artemisinins act at inhibitory synapses in the central nervous system. The present study shows that artemisinins not only act at the postsynaptic side, but also affect the functionality of the presynaptic terminals via their interaction with PDXK which ultimately leads to a decrease in neurotransmitter biosynthesis (Fig. 6). Although the data presented here and our earlier study (14) define mechanisms underlying the downregulation of inhibitory neurotransmission by artemisinins, other neurotransmitters such as dopamine, histamine, and serotonin are also synthesized in a PLP-dependent manner (4–6); thus, future studies will be required to comprehensively dissect the molecular details underlying the artemisinin-induced regulation of neurotransmitter levels and the resulting physiological consequences.

In addition, a comparison of the artemisinin binding pockets in mPDXK and GephE revealed common denominators of drug recognition. Notably, in both cases, artemisinins engage in crucial van der Waals interactions with aromatic residues. In addition, in GephE as well as PDXK, the side chain of an Arg, contributes to artesunate binding, which stabilizes the drug through an electrostatic interaction with its succinate moiety, thus revealing common signatures of artemisinin binding pockets. Our results thus not only broaden the understanding of target recognition by artemisinins at the structural level, but also provide important insights into how these interactions impair inhibitory synaptic transmission in the brain and how this might

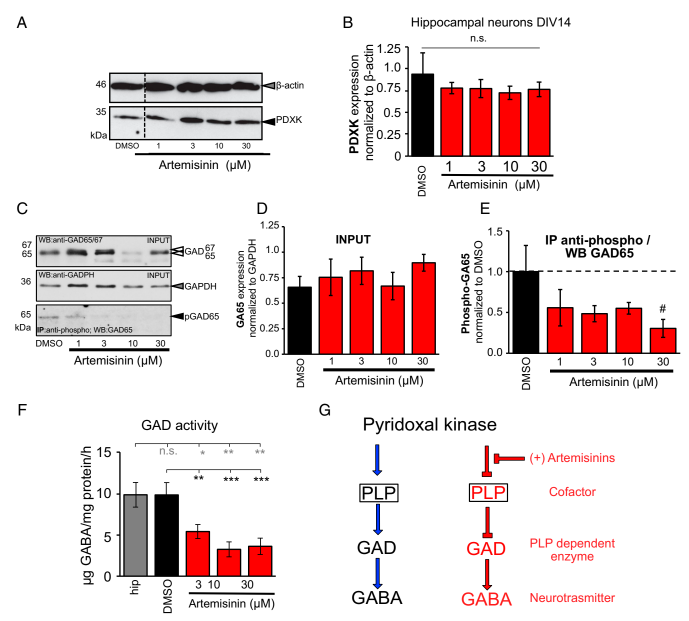


Fig. 5. Artemisinins impact GABA biosynthesis by down-regulating GAD activity. (A) Representative image of a Western blot stained for PDXK (35 kDa, black arrowhead) and β -actin (46 kDa, gray arrowhead). DMSO-treated cells served as internal control. (B) PDXK expression analysis after incubation with artemisinins (1, 3, 10, and 30 µM). PDXK expression was normalized to β -actin. (C) Representative images of input probes after artemisinin treatment (1, 3, 10, and 30 µM) used for immunoprecipitation stained for GAD65/67 (65/67 kDa, white arrowheads) and GAPDH (36 kDa, gray arrowhead). Following immunoprecipitation using an anti-phospho antibody, probes were stained with a specific GAD65 antibody (65 kDa, black arrowhead). DMSO-treated cells served as control samples. (D) Quantitative analysis of input probes from lysates of hippocampal neurons DIV14 incubated with increasing concentrations of artemisinin (1, 3, 10, and 30 µM; DMSO served as control) normalized to the GAPDH signal. (E) Precipitated pGAD (phosphorylated GAD65) was normalized to the DMSO control. Note, that the number of experiments was ${}^{\#}n = 2$ for 30 μ M artemisinin; all other data result from n = 4. All values are shown as mean \pm SEM. (F) Measurements of GAD activity in hippocampal samples following treatment with different concentrations of artemisinin (3, 10, and 30 μM). Tissues without treatment (gray bar) and treated with DMSO (black bar) served as positive controls. Number of measurements n = 8 to 9 from three independent biological replicates. GAD activity decreased significantly with increasing concentrations of artemisinin. The data were analyzed with a paired t test. **P = 0.0054, ***P = 0.0007, and ***P = 0.0002 against DMSO measurements and *P = 0.018, **P = 0.0087, and **P = 0.0047 against hippocampal measurements for artemisinin concentrations of 3, 10, and 30 µM. (G) Schematic representation of the steps leading to GABA biosynthesis at presynaptic terminals. Left shows how GAD synthesizes GABA by utilizing PLP as a cofactor which is produced by PDXK, while Right shows how artemisinins inhibit the initial step in the biosynthesis by inhibiting PDXK, which, in turn, indirectly impacts downstream biosynthetic processes and eventually down-regulates the amount of neurotransmitter being synthesized.

account for the neurological side effects of these drugs. Future studies, along with the molecular signatures revealed in our structures, will be required to investigate whether artemisinins indeed directly bind to and also modulate the activities of other mammalian targets such as protein disulfide isomerase and fatty acid synthase which were identified earlier (7).

Materials and Methods

Experimental Model and Subject Details. For cloning purposes *Escherichia coli* DH5 α was used and the cells were grown on Luria Broth (LB)-agar plates and in LB liquid medium at 37 °C. For recombinant protein expression *E. coli* SoluBL21 cells were used. The cells were grown at 37 °C initially and were further incubated at 30 °C for 16 to 18 h after induction. Transverse

BIOCHEMISTRY

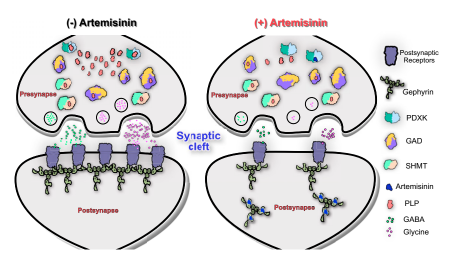


Fig. 6. Schematic representation of inhibitory synapses in the absence and presence of artemisinins. This scheme shows that in the absence of artemisinins (*Left*) gephyrin clusters the receptors required for inhibitory neurotransmitters at postsynaptic sides, while PDXK contributes to the biosynthesis of neurotransmitters at presynaptic terminals by producing the PLP cofactor for GAD and SHMT enzymes. In contrast, in the presence of artemisinins (*Right*), gephyrin-mediated clustering of receptors at postsynaptic sites is impaired while neurotransmitter biosynthesis at presynaptic terminals is inhibited.

hippocampal slices (350 µm thick) were prepared from sevofluraneanesthetized adult C57BL/6J mice (2 to 4 months old) of either sex purchased from Charles River. Adult animals (12-week-old male or female mice) were taken from the mouse strain CD1 (strain code 022; Charles River) to isolate the hippocampi. Animals were housed under standard conditions and all procedures were conducted according to the guidelines and with approval of the local government of Lower Franconia. Preparation of brain slices containing the hippocampal formation was performed as previously described (38).

Method Detail.

Cloning, recombinant protein expression, and purification. The cDNA encoding mPDXK was subcloned into the pETM14 expression vector harboring a 3C precision protease cleavage and BamH1 sites by sequence-independent ligation cloning (SLIC) (39). The proteins (WT and all mutants) were expressed in the E. coli SoluBL21 strain. Cells were grown at 37 °C and expression was induced with 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at an optical density (OD₆₀₀) of 0.6 to 0.8 and cultures were subsequently incubated at 30 °C for 16 to 18 hours. Following centrifugation at 8,000 × g for 15 min the harvested cells were resuspended in lysis buffer containing 50 mM Tris pH 8, 300 mM NaCl, and 5 mM β-mercaptoethanol (β-ME) and lysis was performed by using a microfluidizer. For purification, a two-step protocol was employed consisting of an initial Ni-affinity chromatography with Ni-iminodiacetic acid (IDA) beads, which was followed by cleavage of the N-terminal His₆ tag by incubation with 3C precision protease overnight at 4 °C. Finally, size exclusion chromatography on a Superdex 200 26/60 (GE Healthcare) column was performed in SEC buffer (20 mM Tris pH 8, 150 mM NaCl, and 5 mM β -ME) to purify the protein to apparent homogeneity.

SEC-MALLS. SEC-MALLS experiments of 100 μM WT and all mutants were carried out by using a Superdex 200 10/300 column (GE Healthcare) in SEC buffer. The experiments were performed at a constant flow rate of 0.5 mL/ min at room temperature. The differential refractive index (dRI) and the light-scattering (LS) signals were monitored with a Dawn Helios detector from Wyatt Technologies and molecular masses were derived from the dRI and LS measurements.

Crystallization. Crystallization of mPDXK was performed in the apo-form and in complex with ATP₇S at a protein concentration of 12 mg/mL corresponding to a molar concentration of 0.3 mM. The protein was mixed with 2 mM ATP₇S and 5 mM MgCl₂ and the complex was incubated on ice for 30 min prior to crystallization. Crystallization was performed with the sitting drop vapor diffusion method by mixing equal volumes of protein and mother liquor at 20 °C. The mPDXK-ATP₇S-artesunate structure was determined by soaking mPDXK-ATP₇S crystals with different concentrations of artesunate (2 to 10 mM) for 30 to 600 s. Crystals were transferred into mother liquor (0.18 to 0.24 M sodium thiocyanate and 18 to 26% PEG3350) supplemented with different concentrations of artesunate and 25% glycerol as cryoprotectant before flash cooling in liquid nitrogen.

Data collection, structure determination, and refinement. Data collection for all crystals was performed at the European Synchrotron Radiation Facility (ESRF),

Grenoble, France, on beamline ID23A-1 at a wavelength of 0.9724 Å at 100 K. Datasets were indexed and integrated with XDS (40) and subsequently scaled and merged with AIMLESS (41) from the CCP4 suite (42). The apostructure and the binary mPDXK-ATP γ S complex were determined by molecular replacement with PhaserMR (43) using the human PDXK structure (PDB: 2YXT) as search model and the ternary mPDXK-ATP γ S-artesunate complex was solved with the apo-mPDXK structure as search model. The protein crystallized in space group C2 with four molecules in the asymmetric unit. Refinement was performed in PHENIX (44) with repeated manual model building in Coot (45). Coordinates and restraints for artesunate were obtained from our gephyrin–artesunate structure (PDB: 6FGC). All figures representing protein structures were generated with PyMOL (Schrodinger LLC), Chimera (46), or ChimeraX (47).

Enzymatic activity assay. Pyridoxal kinase activity (WT and variants) was measured following a previously described procedure (22). Briefly, the assay was conducted in 10 mM Hepes buffer (pH 7.3) at 37 °C with 100 mM KCl, 1 mM MgCl₂, 1 mM Mg-ATP, and 50 µg/mL BSA. The pyridoxal kinase concentration was 20 μ g/mL (0.6 μ M), and the substrate pyridoxal was added in a range from 10 to 600 µM. The activity was measured following the increase in absorbance at 388 nm due to PLP formation (extinction coefficient of 4,900 M⁻¹·cm⁻¹) in a CLARIOstar (BMG LABTECH) microplate reader. All experiments were carried out in triplicate. K_M and k_{cat} values were calculated by a Lineweaver-Burk plot (48) with the program Prism (GraphPad Software). For statistical significance of the enzymatic assays, initially, the normality distribution of the data was determined by a D'Agostino and Pearson normality test. After passing the normality test, the statistical significance was determined by a paired t test. For all statistical tests, the P values correspond to *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001; ns, not significant. Statistical analyses were performed by using values from four independent experiments.

To derive the K_i values, the assay was performed under the same conditions using pyridoxal at concentrations of 50 and 150 μ M. Using both pyridoxal concentrations the assays were performed with twofold serial dilutions of artesunate and artemisinin, starting at concentrations of 2.5 and 0.156 mM, respectively. K_i values for the inhibitors artesunate and artemisinin were estimated by a Dixon plot (49), by using a linear regression fit (*P* < 0.001) of the inverted velocity values. The K_i value corresponds to the intersection between the two lines obtained for each individual pyridoxal concentration. For determining the IC₅₀ values, the values of inhibitor concentration were transformed to a logarithmic scale and fitted using a non-linear regression fit with variable slope. IC₅₀ values were calculated as the concentration of inhibitor that gives a velocity halfway between the minimal and maximal values of the curve. All curve fitting procedures and statistical analyses were performed using Prism (GraphPad Software).

Electrophysiology. Transverse hippocampal slices (350 μ m thick) were prepared from adult C57BL/6J mice. Animals were housed under standard conditions and all procedures were conducted according to the guidelines and with approval of the local government of Lower Franconia. Whole-cell voltage-clamp recordings were obtained from visualized pyramidal cells of the

hippocampal CA1 region in a submerged chamber with perfusion solution containing 125 mM NaCl, 3 mM KCl, 2.5 mM CaCl₂, 1.5 mM MgCl₂, 1.25 mM NaH₂PO₄, 25 mM NaHCO₃, and 10 mM D-glucose at 31 °C, constantly gassed with 95% O₂ / 5% CO₂ (pH 7.4). mIPSCs were recorded in the presence of the ionotropic glutamate receptor antagonist kynurenic acid (2 mM) and tetrodotoxin (TTX) (0.5 μ M) using pipettes filled with solution containing 130 mM CsCl, 5 mM Hepes, 3 mM MgCl₂, 5 mM EGTA, 2 mM Na₂ATP, 0.3 mM Na₃GTP, 4 mM NaCl, and 5 mM QX-314 (pH 7.3). mIPSCs were recorded at a holding potential of -70 mV as downward deflections. Current signals were filtered at 2 kHz and sampled at 20 kHz using a Multiclamp 700B amplifier together with a Digidata 1440A interface and the pClamp10 software (Molecular Devices). Data were analyzed off line with Clampfit 10.6 (Molecular Devices) and were expressed as means \pm SEM.

Statistical comparisons of drug effects were performed with OriginPro 2015G (OriginLab Corporation) using a paired t test. Significance was assumed for P values <0.05.

Fluorimetric assay for GAD activity determination. The basis for the determination of the activity of GAD in brain tissue is the ninhydrin reaction. During this reaction, the substrate ninhydrin reacts with the amino group of GABA, releasing water and forming a Schiff base. Following decarboxylation of GABA and elimination of the amino acid, amino-ninhydrin is formed, which dimerizes with ninhydrin (blue color). The concentration of ninhydrin is proportional to the concentration of the amino acid (law of Lambert-Beer) and was measured with a FluoroMax-4 (HORIBA Scientific) fluorimeter using an excitation wavelength of 375 nm (6 mm slide width) and an emission wavelength of 450 nm (10 mm slide width) (50, 51).

The experimental setup was slightly modified from ref. 51. Hippocampi were transferred into a fresh tube containing sonification buffer (0.5 M KCl, 0.01 M EDTA, 0.5% Triton X-100 in sodium-phosphate buffer, pH 6.4) followed by sonification for 10 s. Protein concentration was measured via a Bradford assay and adjusted to 1 $\mu g/\mu L$. The following conditions were analyzed in triplicates: hippocampi without treatment, DMSO, 3 μ M artemisinin, 10 μ M artemisinin, and 30 μ M artemisinin. For each sample a probe (P) and blank (B) were used with 100 μ L homogenized hippocampi and the corresponding DMSO/artemisinin concentrations (3, 10, and 30 μ M) diluted in sonification buffer. The B fraction was supplemented with 200 μ L 10% trichloroacetic acid (TCA). The P and B fractions were supplemented with 100 μ L substrate buffer, pH 6.7, 40 μ L 50 mM pyridoxal phosphate 5-phosphate buffer) and incubated for 2 hours at 38 °C.

Subsequently the reaction of the P fraction was also stopped by adding 200 μ L 10% TCA. Samples were centrifuged at 950 \times g for 20 min. A total of 200 μ L of all probes was incubated with 400 μ L ninhydrin (14 mM in 0.5 M sodium carbonate buffer, pH 9.93) at 60 °C for 30 min. Afterward, probes were incubated with 9 mL copper tartrate (1.6 g sodium carbonate, 329 mg tartaric acid, 300 mg copper-(II) sulfate in 1 L aqua dest) at 22 °C for 20 min and measured (1:10 diluted) in the fluorescence spectrophotometer. A GABA concentration series of 0, 0.5, 1, 3, 5, 20, 50 μ M GABA was used as standard.

Preparation of primary hippocampal neurons. Hippocampal neurons were prepared at embryonic day 17 (E17) from pregnant female wild-type mice. Dissociated cells were grown in neurobasal medium supplemented with 5 mL of L-glutamine (200 mM) and B27 supplement (Life Technologies; A3582801) with an exchange of 50% medium after every 6 d in culture.

Protein lysate preparation. At DIV14, hippocampal neurons were incubated for 2 hours with different DMSO/artemisinin concentrations (1, 3, 10, and 30 μ M). Cells were washed and harvested in phosphate-buffered saline (PBS), pH 7.4, with the help of a cell scraper. After a centrifugation step, the pellet was resuspended in 100 μ L of brain homogenate buffer (20 mM Hepes, 100 mM K-acetate, 40 mM KCl, 5 mM EGTA, 5 mM MgCl₂, 5 mM dithiothreitol (DTT), 1 mM phenylmethylsulfonyl fluoride (PMSF), 1% Triton X, protease inhibitor Roche complete, pH 7.2) and sonicated at low power for 5 s. Protein concentration was used for Western blot analysis.

Western blot. For SDS/PAGE, 11% polyacrylamide gels were freshly prepared, followed by Western blot on nitrocellulose membranes (GE Healthcare).

- 1. M. H. Li *et al.*, Conformational changes in the reaction of pyridoxal kinase. *J. Biol. Chem.* **279**, 17459–17465 (2004).
- J. T. Neary, W. F. Diven, Purification, properties, and a possible mechanism for pyridoxal kinase from bovine brain. J. Biol. Chem. 245, 5585–5593 (1970).
- C. Merigliano, E. Mascolo, R. Burla, I. Saggio, F. Vernì, The relationship between vitamin B6, diabetes and cancer. Front. Genet. 9, 388 (2018).

Membranes were blocked for 1 hour with 5% BSA in TBS-T (TBS with 1% Tween 20). Primary antibodies were incubated overnight at 4 °C. GAD and PDXK proteins were detected with the GAD67/65-specific antibody (ab11070 1:1,000; abcam) and PDXK-specific antibody (NBP1-88283, 1:1,000; novusBio). β -Actin (GTX26276, 1:5,000; GeneTex/Biozol) served as loading control.

Signals were detected using the ECL plus system (GE Healthcare).

Data analysis of Western blots. Image quantification was performed using ImageJ (1.51)/Fiji² (52–54). The data were analyzed using a Student's t test (analysis of variance) and values below *P < 0.05 were considered significant, **P < 0.01, ***P < 0.001. The values are displayed as mean \pm SEM or as otherwise noted.

Immuncytochemical staining. DIV14 primary hippocampal neurons were incubated for 2 hours with corresponding DMSO/artemisinin concentrations (1, 3, 10, and 30 μ M). Neurons were fixed in 4% paraformaldehyde in PBS for 15 min. After washing twice with PBS, 50 mM NH₄Cl was added for 10 min followed by blocking with 5% goat serum in PBS (permeabilized with 0.2% Triton X-100) for 30 min at 22 °C. Primary antibodies were incubated for 1 hour in blocking solution without Triton X-100. GAD and PDXK proteins were detected with the GAD67/65-specific antibody (ab11070, 1:500; abcam), gephyrin-specific antibody (147111, 1:500; Synaptic Systems) and PDXK-specific antibody (NBP1-88283, 1:150; novusBio). Secondary antibodies gamCy3, garAlexa488, and garCy5 (1:500; Dianova) were applied for 1 hour. Cells were stained with 4',6 diamino-2-phenylindole (DAPI) and slides were mounted with Moviol.

Immunoprecipitation. DIV14 primary hippocampal neurons were incubated for 2 hours with corresponding DMSO/artemisinin concentrations (1, 3, 10, and 30 µM). Cells were lysed in lysis buffer (TBS: 50 mM Tris, 150 mM NaCl, pH 8.0) containing 0.1% Triton X-100, 0.1 mM PMSF, 2 mM EDTA, and complete, EDTA-free protease inhibitor mixture (Roche) and sonicated. The protein concentration of each probe was measured, and 50 μ g was used for immunoprecipitation (IP). The protein solution (300 µL) was supplemented with an anti-phospho antibody (1:100, ab17464; abcam) and agitated overnight at 4 °C. A total of 40 µL of protein A-Sepharose beads (REF 11719408001; Roche) was added and agitation continued overnight at 4 °C. Following a short centrifugation step, the supernatant was removed, and beads were washed three times with lysis buffer. Proteins were eluted from beads by incubation with 30 μ L of 2× SDS sample buffer at 95 °C for 7 min. Proteins were separated on 11% SDS gels and stained after Western blotting with GAD65 antibody (1:1,000, ab26113; abcam). Input controls were stained with GAPDH antibody (1:1,000, CB1001; Calbiochem) and GAD65/67 antibody (1:1,000, ab11070). All experiments were performed with at least three independent biological replicates (unless otherwise stated). Quantification of input GAD65 protein signals at different artemisinin concentrations (1, 3, 10, and 30 μ M) were normalized to GAPDH signals of corresponding input probes. The IP probes were finally normalized to the DMSO signal. Calculated data were compared using an unpaired t test with a probability of error of P < 0.05 considered to be significant.

Data Availability. The coordinates of mPDXK-apo, mPDXK-ATP_YS, and mPDXK-ATP_YS-artesunate structures have been deposited in the PDB with accession codes 6YJZ, 6YK0, and 6YK1, respectively.

ACKNOWLEDGMENTS. We thank Dr. Antje Gohla for providing the cDNA of mPDXK and Dana Wegmann and Christine Schmitt for excellent technical assistance. We also thank the beamline scientists at beamline ID23-A1, ESRF, Grenoble, for technical assistance during data collection. We are grateful to Dr. Kunimichi Suzuki for the critical reading of the manuscript. This work was supported by the Deutsche Forschungsgemeinschaft (SCHI425/8-2) and the Rudolf Virchow Center for Integrative and Translational Bioimaging (H.S.). A.P.M. was supported by the Graduate School of Life Sciences at the University of Würzburg. N.S. was supported by funds of the Bavarian State Ministry of Science and the Arts and the Graduate School of Life Sciences. V.B.K. is supported at the Medical Research Council (MRC) Laboratory of Molecular Biology (LMB) by a European Molecular Biology Organization (EMBO) Long-Term Fellowship (ALTF137-2019).

- M. L. di Salvo, M. K. Safo, R. Contestabile, Biomedical aspects of pyridoxal 5'phosphate availability. Front. Biosci. (Elite Ed.) 4, 897–913 (2012).
- R. Percudani, A. Peracchi, A genomic overview of pyridoxal-phosphate-dependent enzymes. *EMBO Rep.* 4, 850–854 (2003).
- A. C. Eliot, J. F. Kirsch, Pyridoxal phosphate enzymes: Mechanistic, structural, and evolutionary considerations. Annu. Rev. Biochem. 73, 383–415 (2004).

- 7. J. Li et al., Artemisinins target GABAA receptor signaling and impair alpha cell identity. Cell 168, 86–100.e15 (2017).
- Y. Tu, Artemisinin-A gift from traditional Chinese medicine to the world (nobel lecture). Angew. Chem. Int. Ed. Engl. 55, 10210–10226 (2016).
- WHO, "Guidelines for the treatment of malaria" in *Guidelines for the Treatment of Malaria* (World Health Organization, Geneva, Switzerland, ed. 3, 2015).
- M. P. Crespo-Ortiz, M. Q. Wei, Antitumor activity of artemisinin and its derivatives: From a well-known antimalarial agent to a potential anticancer drug. J. Biomed. Biotechnol. 2012, 247597 (2012).
- 11. A. Gautam, T. Ahmed, V. Batra, J. Paliwal, Pharmacokinetics and pharmacodynamics of endoperoxide antimalarials. *Curr. Drug Metab.* **10**, 289–306 (2009).
- T. van der Meulen et al., Artemether does not turn alpha cells into beta cells. Cell Metab. 27, 218–225.e214 (2018).
- A. M. Ackermann, N. G. Moss, K. H. Kaestner, GABA and artesunate do not induce pancreatic alpha-to-beta cell transdifferentiation in vivo. *Cell Metab.* 28, 787–792.e783 (2018).
- V. B. Kasaragod et al., Elucidating the molecular basis for inhibitory neurotransmission regulation by artemisinins. *Neuron* 101, 673–689.e611 (2019).
- V. B. Kasaragod, H. Schindelin, Structural framework for metal incorporation during molybdenum cofactor biosynthesis. *Structure* 24, 782–788 (2016).
- J. Kuper, A. Llamas, H. J. Hecht, R. R. Mendel, G. Schwarz, Structure of the molybdopterin-bound Cnx1G domain links molybdenum and copper metabolism. *Nature* 430, 803–806 (2004).
- V. B. Kasaragod, H. Schindelin, Structure-function relationships of glycine and GABA_A receptors and their interplay with the scaffolding protein gephyrin. *Front. Mol. Neurosci.* **11**, 317 (2018).
- J. A. Kerry, M. Rohde, F. Kwok, Brain pyridoxal kinase. Purification and characterization. *Eur. J. Biochem.* 158, 581–585 (1986).
- P. W. Elsinghorst, M. L. di Salvo, A. Parroni, R. Contestabile, Inhibition of human pyridoxal kinase by 2-acetyl-4-((1R,2S,3R)-1,2,3,4-tetrahydroxybutyl)imidazole (THI). J. Enzyme Inhib. Med. Chem. 30, 336–340 (2015).
- M. C. Hanna, A. J. Turner, E. F. Kirkness, Human pyridoxal kinase. cDNA cloning, expression, and modulation by ligands of the benzodiazepine receptor. J. Biol. Chem. 272, 10756–10760 (1997).
- D. C. Jones, M. S. Alphey, S. Wyllie, A. H. Fairlamb, Chemical, genetic and structural assessment of pyridoxal kinase as a drug target in the African trypanosome. *Mol. Microbiol.* 86, 51–64 (2012).
- F. Kwok, J. E. Churchich, Brain pyridoxal kinase. Purification, substrate specificities, and sensitized photodestruction of an essential histidine. J. Biol. Chem. 254, 6489–6495 (1979).
- D. B. McCormick, M. E. Gregory, E. E. Snell, Pyridoxal phosphokinases. I. Assay, distribution, I. Assay, distribution, purification, and properties. J. Biol. Chem. 236, 2076–2084 (1961).
- M. K. Safo et al., Crystal structure of pyridoxal kinase from the Escherichia coli pdxK gene: Implications for the classification of pyridoxal kinases. J. Bacteriol. 188, 4542–4552 (2006).
- J. B. Ubbink, S. Bissbort, W. J. Vermaak, R. Delport, Inhibition of pyridoxal kinase by methylxanthines. *Enzyme* 43, 72–79 (1990).
- F. N. Musayev et al., Crystal structure of human pyridoxal kinase: Structural basis of M(+) and M(2+) activation. Protein Sci. 16, 2184–2194 (2007).
- M. H. Li et al., Crystal structure of brain pyridoxal kinase, a novel member of the ribokinase superfamily. J. Biol. Chem. 277, 46385–46390 (2002).
- D. Engel et al., Plasticity of rat central inhibitory synapses through GABA metabolism. J. Physiol. 535, 473–482 (2001).

- J. Wei, K. M. Davis, H. Wu, J. Y. Wu, Protein phosphorylation of human brain glutamic acid decarboxylase (GAD)65 and GAD67 and its physiological implications. *Biochemistry* 43, 6182–6189 (2004).
- C. C. Chou et al., Activation of brain L-glutamate decarboxylase 65 isoform (GAD65) by phosphorylation at threonine 95 (T95). Mol. Neurobiol. 54, 866–873 (2017).
- T. M. E. Davis et al., Penetration of dihydroartemisinin into cerebrospinal fluid after administration of intravenous artesunate in severe falciparum malaria. Antimicrob. Agents Chemother. 47, 368–370 (2003).
- G. Schmuck, E. Roehrdanz, R. K. Haynes, R. Kahl, Neurotoxic mode of action of artemisinin. *Antimicrob. Agents Chemother.* 46, 821–827 (2002).
- T. G. Brewer et al., Fatal neurotoxicity of arteether and artemether. Am. J. Trop. Med. Hyg. 51, 251–259 (1994).
- D. L. Wesche, M. A. DeCoster, F. C. Tortella, T. G. Brewer, Neurotoxicity of artemisinin analogs in vitro. Antimicrob. Agents Chemother. 38, 1813–1819 (1994).
- 35. L. Tang et al., Crystal structure of pyridoxal kinase in complex with roscovitine and derivatives. J. Biol. Chem. 280, 31220–31229 (2005).
- S. Bach et al., Roscovitine targets, protein kinases and pyridoxal kinase. J. Biol. Chem. 280, 31208–31219 (2005).
- A. K. Gandhi et al., Crystal structures of human pyridoxal kinase in complex with the neurotoxins, ginkgotoxin and theophylline: Insights into pyridoxal kinase inhibition. PLoS One 7, e40954 (2012).
- F. Zheng et al., Activin controls ethanol potentiation of inhibitory synaptic transmission through GABAA receptors and concomitant behavioral sedation. Neuropsychopharmacology 41, 2024–2033 (2016).
- M. Z. Li, S. J. Elledge, Harnessing homologous recombination in vitro to generate recombinant DNA via SLIC. Nat. Methods 4, 251–256 (2007).
- 40. W. Kabsch, Xds. Acta Crystallogr. D Biol. Crystallogr. 66, 125-132 (2010).
- P. R. Evans, G. N. Murshudov, How good are my data and what is the resolution? Acta Crystallogr. D Biol. Crystallogr. 69, 1204–1214 (2013).
- M. D. Winn et al., Overview of the CCP4 suite and current developments. Acta Crystallogr. D Biol. Crystallogr. 67, 235–242 (2011).
- A. J. McCoy et al., Phaser crystallographic software. J. Appl. Cryst. 40, 658–674 (2007).
 P. D. Adams et al., PHENIX: A comprehensive python-based system for macromolec-
- H. F. D. Adams et al., Hindric A completenensite pythol-based system of indefinition ular structure solution. Acta Crystallogr. Dol. Crystallogr. 66, 213–221 (2010).
- P. Emsley, K. Cowtan, Coot: Model-building tools for molecular graphics. Acta Crystallogr. D Biol. Crystallogr. 60, 2126–2132 (2004).
- E. F. Pettersen et al., UCSF Chimera–A visualization system for exploratory research and analysis. J. Comput. Chem. 25, 1605–1612 (2004).
- T. D. Goddard et al., UCSF ChimeraX: Meeting modern challenges in visualization and analysis. Protein Sci. 27, 14–25 (2018).
- H. Lineweaver, D. Burk, The determination of enzyme dissociation constants. J. Am. Chem. Soc. 56, 658–666 (1934).
- M. Dixon, The determination of enzyme inhibitor constants. *Biochem. J.* 55, 170–171 (1953).
- I. P. Lowe, E. Robins, G. S. Eyerman, The fluorometric measurement of glutamic decarboxylase and its distribution in brain. J. Neurochem. 3, 8–18 (1958).
- M. R. Holdiness, J. B. Justice, B. Darryl, B. D. Neill, J. D. Salamone, Fluorimetric assay for the determination of glutamic acid decarboxylase activity in subregions of rat brain tissue. *Anal. Lett.* **13**, 1333–1344 (1980).
- C. A. Schneider, W. S. Rasband, K. W. Eliceiri, NIH image to ImageJ: 25 years of image analysis. Nat. Methods 9, 671–675 (2012).
- J. Schindelin, C. T. Rueden, M. C. Hiner, K. W. Eliceiri, The ImageJ ecosystem: An open platform for biomedical image analysis. *Mol. Reprod. Dev.* 82, 518–529 (2015).
- J. Schindelin et al., Fiji: An open-source platform for biological-image analysis. Nat. Methods 9, 676–682 (2012).

BIOCHEMISTRY