Modulation of Voltage-dependent Shaker Family Potassium Channels by an Aldo-Keto Reductase**

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Jun Weng,1 Yu Cao,1 Noah Moss, and Ming Zhou2

From the Department of Physiology and Cellular Biophysics, College of Physicians and Surgeons, Columbia University, New York, New York 10032

The β subunit (Kvβ) of the Shaker family voltage-dependent potassium channels (Kv1) is a cytosolic protein that forms a permanent complex with the channel. Sequence and structural conservation indicates that Kvβ resembles an aldo-keto reductase (AKR), an enzyme that catalyzes a redox reaction using an NADPH cofactor. A putative AKR in complex with a Kv channel has led to the hypothesis that intracellular redox potential may dynamically influence the excitability of a cell through Kvβ. Since the AKR function of Kvβ has never been demonstrated, a direct functional coupling between the two has not been established. We report here the identification of Kvβ substrates and the demonstration that Kvβ is a functional AKR. We have also found that channel function is modulated when the Kvβ-bound NADPH is oxidized. Further studies of the enzymatic properties of Kvβ seem to favor the role of Kvβ as a redox sensor. These results suggest that Kvβ may couple the excitability of the cell to its metabolic state and present a new avenue of research that may lead to understanding of the physiological functions of Kvβ.

Kv channels control the flow of K⁺ through the cell membrane in response to changes in membrane potential. Opening of the channel causes membrane hyperpolarization that can curtail excessive membrane excitability or simply tone down normal membrane activity. Kv channels control the flow of K⁺ through the cell membrane in response to changes in membrane potential. Opening of the channel causes membrane hyperpolarization that can curtail excessive membrane excitability or simply tone down normal membrane activity. Kvβ, found in plants, insects, and mammals, attaches to the cytosolic face of the Kv1 channels to form a macromolecular complex (Fig. 1A) (1–5). There are three mammalian Kvβ genes (Kvβ1–3) (6, 7), and all Kvβs have a conserved core region with sequence similarities to an AKR (8, 9). The structure of the conserved core of a Kvβ, Kvβ2 from rat, shows that it has a canonical AKR fold, with highly conserved catalytic residues in the correct geometry for catalysis to happen, and has a nicotinamide cofactor tightly bound (Fig. 1B) (3, 10, 11). These features suggest that Kvβ is a functional AKR. In many cells, acute changes in Kv channel current have long been observed when the cellular redox state is altered, for example, under hypoxic condition or oxidative stress (12–17), but the redox-sensing mechanism remains unknown. Could Kvβ be the missing link between cellular redox chemistry and Kv channel activities? In this study, we address the questions of whether Kvβ is a functional AKR and whether the AKR function is coupled to channel functions.

AKRs catalyze the reduction of an aldehyde to an alcohol by oxidizing an NADPH cofactor. Most AKRs have a broad substrate spectrum so that in addition to their native substrates, they also convert small molecule aldehydes such as benzaldehyde derivatives (18). The catalysis follows a kinetic mechanism shown in Scheme I. An AKR, E, binds sequentially an NADPH and an aldehyde substrate (19) to form a ternary complex, E-NADPH-aldehyde. The redox reaction then occurs when the enzyme helps transfer a hydride from the cofactor to the aldehyde. The product, an alcohol, dissociates from the enzyme and is followed by the oxidized cofactor NADP⁺. Many AKRs show high specificity to NADPH over its close relative, NADH (18). The hydride transfer step is usually fast, and the nucleotide exchange steps, or the protein conformational changes associated with them, are slow and rate-limiting (20, 21). The cycle in Scheme I is reversible so that an alcohol can be oxidized to an aldehyde accompanied by the conversion of an NADP⁺ to NADPH.

EXPERIMENTAL PROCEDURES

Molecular Biology and Mutagenesis—For protein expression, rat Kvβ2 (GenBank™ accession number: CAA54142, residues 36–363) was cloned into a pQE70 vector (Qiagen) between the SpII and BamHI sites with a C terminus His8 tag. For electrophysiological measurements, rat Kv1.4 (GenBank™ accession number: CAA34133) or Kvβ2 was cloned into a modified pBluescript vector for in vitro transcription. Mutations were made using the QuikChange (Strategene) kit and verified by sequencing through the entire coding region.

Protein Expression and Purification—XL-1Blue cells were used to express protein. Cells were grown in Luria broth at 37 °C to an OD of ~1.2 and induced with 0.5 mM isopropyl β-D-thiogalactopyranoside (final concentration). Immediately after induction, the temperature was reduced to 20 °C, and cells were harvested 16 h after induction. Expressed protein was purified on a Talon Co2⁺ affinity column (BD Biosciences). Throughout the purification procedure, the following buffer was used: 20 mM Tris, pH 8.0, 300 mM KCl, 10% (v/v) glycerol, and 1 mM β-mercaptoethanol. Non-specifically bound protein was washed using 20 mM imidazole added to the above solution, and Kvβ2 protein was eluted with 300 mM imidazole. Immediately after elution, the affinity tag was removed by a brief incubation with trypsin at a Kvβ2 to trypsin ratio of ~150:1 (w/w). The protein was then loaded onto a Superdex 200 column (Amersham Biosciences) for final purification. The column was equilibrated with the reaction buffer: 150 mM KCl, 20 mM Tris, pH 8.0. Protein concentration was determined using the BCA kit (Pierce).

Enzyme Kinetics Measurement—To measure the steady-state multiple turnover reaction rate, the reaction mixture (100 μl) contains 0.2 mM NADPH, 30 μg Kvβ2 protein, and desired concentration of sub-
strates in the reaction buffer. The reaction mixture was incubated at 37 °C and NADPH (ε max = 6.22 mM−1 cm−1) absorption was monitored at 340 nm using a Pharmaspec UV-1700 (Shimadzu) equipped with a temperature controlling unit (CPS-240A). The reaction was initiated by adding the enzyme and monitored the absorbance at 340 nm using a PharmaSpec UV-1700 (Shimadzu) equipped with a temperature controlling unit (CPS-240A). The reaction was initiated by the addition of a substrate. The initial turnover rate constant was calculated from the linear part of the decrease in absorbance. Blank controls with no protein were incorporated routinely, and the background NADPH consumption was subtracted.

As a positive control, we obtained rat 3α-hydroxysteroid dehydrogenase protein (a gift from Dr. Trevor Penning, University of Pennsylvania) and measured its specific activity using the following conditions: 75 μM androsterone (dissolved in acetonitrile), 2.3 mM NAD⁺, 100 mM potassium phosphate, pH 7.0, and 4% acetonitrile (total). We initiated the reaction by adding the enzyme and monitored the absorbance at 340 nm over time. We obtained a specific activity of 1.4 ± 0.1 μM (n = 5) of androsterone oxidized per minute per milligram of protein, similar to the value of 1.6 μM min⁻¹ mg⁻¹ obtained in Dr Penning’s laboratory.

To measure the single turnover hydride transfer reaction rate, Kvβ2 protein was concentrated to 3–10 mg/ml and mixed with proper concentrations of substrates, and the absorption at 363 nm was followed over time at 37 °C. The 23 nm red shift of the absorption peak for the bound NADPH is likely caused by interactions between the nicotinamide ring and the surrounding residues. Previous studies have demonstrated that the 363 nm peak is due to the bound NADPH (22). The fraction of NADPH remaining on Kvβ was then plotted versus time, and the data points were fit with a single exponential function in Origin (Microcal Inc.). The inverse of the exponential time constant is defined as the hydride transfer rate constant. When higher than 5 mM 4-carboxybenzaldehyde (4-CB) was used in either the multiple or single turnover measurement, 40 mM Tris was used to maintain the solution pH at 8.0.

Channel Expression and Electrophysiology—mRNA was prepared by in vitro transcription and purified using the Trizol reagent (Invitrogen). mRNAs were injected into Xenopus oocytes for channel expression. For co-expression, Kv1.4 and Kvβ2 mRNAs were injected at a ratio of 1:3 (w/w). The wild type Kv1.4 has a cysteine residue at position 13 that can be oxidized on inside-out patches to affect channel inactivation, and a C13S mutant or addition of dithiothreitol abolishes this effect (23). To eliminate concerns that the change in inactivation could be due to cysteine oxidation, we used the C13S Kv1.4 in this study. In parallel, we also tested 4-cyanobenzaldehyde (4-CY) and 4-CB on Kv1.4 wild type co-expressed with Kvβ2, in the presence of 5 mM dithiothreitol, and essentially the same results were obtained.

We recorded patch clamp currents from oocytes 3–5 days after injection. Electrodes were drawn from patch glasses (G85150T-4, Warner Instruments) and polished (MP-803, Narishige Co.) to a resistance of 0.6–1 MΩ. The pipette solution contained (in mM): 130 KCl, 2 MgCl₂, and 10 KH₂PO₄ at pH 7.4. The bath solution contained (in mM): 80 KCl, 5 EGTA, and 50 KH₂PO₄ at pH 7.4. pH was adjusted with KOH. The higher buffer content in the bath solution is necessary to maintain the pH at 7.4 for 5 mM or higher 4-CB. K⁺ currents were elicited by holding the patch at −100 mV for at least 30 s and stepping to +60 mV for 500 ms. The volume of the recording chamber (Warner Instruments) is ~120 μl, and complete exchange of solution was achieved by perfusing 1 ml of new solution by gravity flow with a flow rate of 2–3 ml/min.

Inactivation of Kv1.4 with or without Kvβ2 has been studied extensively (24–28). Current inactivation can be fit by an exponential function with two components. The faster component, which is also the predominant one, is mainly contributed by the N-type inactivation (29), and the slower component is thought to be mainly due to the C-type inactivation (27, 30). Here we follow the same tradition to quantify channel inactivation. For Kv1.4 co-expressed with the wild type Kvβ2, both the slow and fast inactivation time constants were modulated by 4-CY (Table 1). Since the N- and C-type inactivations are tightly coupled, we do not know yet if the change in the slow component is due to a direct effect on the C-type inactivation or a result of change in the N-type inactivation. We focused on the fast component, and we defined the inverse of the smaller time constant as the rate constant of channel inactivation and used the rate constant as a measure for the effect of Kvβ substrates.

Chemical Reagents—All chemical reagents were purchased from Sigma, except for 4-oxonononalen (Cayman Chemicals). Substrates were...
first prepared in ethanol (4-CY) or dimethyl sulfoxide (4-CB) and then diluted to the desired final concentration. The final concentration of ethanol or dimethyl sulfoxide is less than 1% in electrophysiological measurements. NADPH was purchased as a tetrasodium salt. Since sodium ions block potassium channels from the intracellular side, we exchanged the sodium to potassium ions by fast protein liquid chromatography, and the stock solution was aliquoted and stored at \(-80^\circ C\).

RESULTS

Single Turnover Enzymatic Reactions—Freshly purified Kv\(\beta_2\) protein contains the reduced form of the cofactor (NADPH), as indicated by a 363 nm peak in the UV absorption spectrum (Fig. 2A), and the occupancy of NADPH is more than 90% (10, 22). Since NADP\(^+\) has little absorption at the same wavelength, the NADPH-to-NADP\(^+\) conversion eliminates the 363 nm peak. Utilizing this as a readout, we screened small molecule aldehydes, which are common AKR substrates. We have identified several potential Kv\(\beta_2\) substrates and two of the compounds, 4-CB and 4-CY (Fig. 2B, inset), were used here to study the enzymatic activity of Kv\(\beta_2\) and substrate-induced channel modulations.

Shown in Fig. 2B are UV spectra recorded at the indicated time points after mixing Kv\(\beta_2\) with 5 mM 4-CB. The peak at 363 nm decreases over time, and the reduction of the peak reflects the oxidation of the Kv\(\beta_2\)-bound NADPH. Since no free NADPH was added, this is a single turnover reaction. When the fraction of NADPH remaining was

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**FIGURE 2.** Enzymatic properties of Kv\(\beta_2\). A, a UV spectrum of freshly purified Kv\(\beta_2\) protein. The peak at 363 nm is magnified in the inset. B, UV spectra taken at the marked time point after mixing Kv\(\beta_2\) protein with 5 mM 4-CB. The chemical structures of 4-CB (R = COOH) and 4-CY (R = CN) are shown in the inset. C, fraction of the 363 nm peak remaining plotted versus time for the wild type (4-CB (C)) and 4-CY (□)) and the D85N mutant (○) Kv\(\beta_2\). The smooth curves are single component exponential functions fit to the data. Error bars are S.E. of five (wild type, 4-CB), six (wild type, 4-CY), and three (D85N) independent experiments. D, UV spectra taken at 0 (black trace), 30 (red trace), and 60 (blue trace) minute(s) after mixing 5 mM 4-CB with NADPH. E, gel filtration elution profiles for oxidized Kv\(\beta_2\) (black trace) and for NADP\(^+\) alone (red trace). The molar ratio of Kv\(\beta_2\) to NADP\(^+\) is kept at 1:1. UV absorption was monitored at 254 nm where NADP\(^+\) has prominent absorption. F, fraction of the total 363 nm peak plotted versus time after the oxidized Kv\(\beta_2\) was mixed with 10 mM 4-MOB. Inset, UV spectra of oxidized Kv\(\beta_2\) taken at the different time point after mixing with 4-MOB. The chemical structure of 4-MOB is shown to the right.
plotted versus time (Fig. 2C), the data points were well fit by a single exponential function with a time constant of 8.4 ± 0.1 min (Fig. 2C), consistent with a single step hydride transfer reaction. Similar results were obtained when 5 mM 4-CY was used but with a smaller time constant of 2.3 ± 0.05 min (Fig. 2C), indicating that 4-CY at 5 mM concentration consumes the bound cofactor ~3-fold faster than 4-CB.

To support the conclusion that the observed NADPH consumption is due to an enzymatic reaction, three control experiments were done. First, free NADPH was mixed with 5 mM 4-CB in the absence of Kvβ2 protein, and the change in NADPH absorption was monitored over time. A very slight decrease of NADPH absorption was observed over a period of 1 h (Fig. 2D), indicating that the reaction is greatly facilitated when the cofactor binds to the Kvβ2 protein.

Second, an active site residue, aspartate 85 (Fig. 1B), was mutated to an asparagine (Asn) and the D85N mutant protein was expressed and purified. Asp85 is highly conserved in all AKRs, and a Asp-to-Asn mutation reduces the rate of catalysis in other AKRs such as aldose reductase and 3α-hydroxysteroid dehydrogenase (3α-HSD) by ~12- and 31-fold, respectively (31, 32). The purified D85N Kvβ2 protein has a bound NADPH cofactor, as indicated by an absorption peak near 360 nm. When the D85N mutant protein was mixed with 5 mM 4-CB, the absorption of the bound cofactor changed very slowly with a time constant of 330 ± 20 min (Fig. 2C; also see Fig. 3A). The time constant is ~40-fold slower than that of the wild type Kvβ, indicating that the mutation slowed down the enzymatic reaction. This result supports the conclusion that Kvβ is a functional AKR.

Third, the reverse reaction, that is, transfer of a hydride from an alcohol to an NADP+, was tested. To do this, we purified the oxidized Kvβ2, i.e. Kvβ2 protein after its bound NADPH was consumed by 4-CY, by passing the reaction mixture through a size exclusion column. Interestingly, we found that the elution does not contain a peak corresponding to NADP+, which should be easily detected (Fig. 2E). This indicates that the oxidized cofactor (NADP+) stays tightly bound to Kvβ2. Using the oxidized Kvβ2, we screened small molecule alcohols for their ability to reduce the bound cofactor, and we found potential substrates. The result from one of them, 4-methoxybenzalcohol (4-MOB), is shown in Fig. 2F. When 4-MOB (10 mM) was mixed with the oxidized Kvβ2, absorption at 363 nm gradually increased (Fig. 2F, inset) and recovers to approximately the same level as freshly purified protein. The time course of NADPH generation was well fit with a single exponential function with a time constant of 61 ± 1 min (Fig. 2F), consistent with a single step hydride transfer reaction. The consumption and the regeneration of the Kvβ2-bound NADPH by small molecule aldehydes and alcohols clearly demonstrated that Kvβ is a functional AKR.

Single Versus Multiple Turnover Enzymatic Reactions—To further characterize the hydride transfer reaction, we measured the rate constant of the bound NADPH consumption, i.e. the single turnover reaction, at different 4-CY or 4-CB concentrations. We found that the rate constant increases with either 4-CB or 4-CY concentration and does not reach saturation even at 20 mM of a substrate (Fig. 3A, left panel). As a control, we used the D85N Kvβ2 protein and measured its hydride transfer rate at three different 4-CB concentrations (Fig. 3A, right panel). The rate is much slower than that of the wild type at all three substrate concentrations (2, 5, and 10 mM), supporting the conclusion that that Kvβ is a functional AKR.

It is remarkable that the rate constant does not saturate with increasing substrate concentrations. This indicates that both substrates have low occupancies on the enzyme so that binding is not saturated, something that is expected because both compounds are not native substrates. More importantly, it indicates that Kvβ is capable of converting the bound NADPH at a rate equal to or faster than 1.9 ± 0.06 min⁻¹ (20 mM 4-CY), a time scale that is physiologically relevant in terms of redox sensing in a cell (33).

The hydride transfer step is only a half-reaction in the enzymatic reaction cycle, so we next measured the steady-state turnover rate constant for the cycle shown in Scheme 1. The turnover rate constant was plotted versus 4-CB or 4-CY concentrations (Fig. 3B), and the data points were well fit by a Michaelis-Menten equation with a maximum turnover rate constant of 0.073 ± 0.0016 and 0.082 ± 0.0011/min and a Km of 3.2 ± 0.17 and 2.7 ± 0.12 mM, for 4-CB and 4-CY, respectively. When NADPH was substituted with NADH, no turnover was observed (data not shown), indicating that Kvβ has evolved to use specifically NADPH for catalysis. The steady-state turnover rate constant for both 4-CB and 4-CY is approximately the same but is much slower than the hydride transfer rate constant, especially for 4-CY. This is consistent with a reaction mechanism in which the cofactor exchange steps are rate-limiting. Since the cofactor exchange steps are not substrate-dependent, it suggests that the steady-state turnover rate constant is similar even when Kvβ catalyzes its native substrate(s). This property may have important physiological implications (see “Discussion”).

As an additional control and to further support the conclusion that Kvβ is an AKR, we mutated another highly conserved catalytic site residue, Lys118 (Fig. 1B), to a methionine. A Lys-to-Met mutation in rat 3α-HSD reduces the multiple turnover rate by more than 1000-fold (32). We expressed and purified the K118M mutant Kvβ2 and measured the multiple turnover rate constant at three different 4-CY concentrations (Fig. 3B). At 5 mM 4-CY, the turnover rate constant ([(1.04 ± 0.21) × 10⁻⁴ min⁻¹] was ~500-fold slower than that of the wild type (0.053 ± 0.003 min⁻¹). The rate is essentially indistinguishable from that of free NADPH oxidation without the presence of a Kvβ protein. This result indicates that the conserved lysine is important for catalysis, and reinforces the conclusion that Kvβ is a functional AKR.
**Redox Modulation of Kv Channels**

**FIGURE 4.** Channel modulations by Kvβ substrates. A–D, current traces of Kv1.4 expressed with no Kvβ2 (B), with the wild type Kvβ2 (A), with the D85N Kvβ2 (C), and with the K118M Kvβ2 (D). In each case, current traces at two different time scales are shown. The black trace was recorded before 4-CY application. After incubating the patch for 10 min in the internal solution containing 5 μM 4-CY, the solution was washed away (black), and the red trace was recorded. E, channel inactivation rate before (black) and after (red) 4-CY application. The error bars are S.E. of 31 (for 1.4 + Kvβ2), 10 (for Kv1.4 alone), 9 (for Kv1.4 + D85N Kvβ2), and 6 (for Kv1.4 + K118M Kvβ2) independent measurements (patches). F, the same experiments shown in A and B. After recording the red trace, the bath solution was exchanged to the one containing 0.2 mM NADPH, and the blue trace was recorded 30 min after incubation. Left panel, Kv1.4 + Kvβ2; right panel, Kv1.4 only. Potassium currents in both Figs. 4 and 5 were recorded from inside-out patches. Currents were elicited by holding the patches at ~100 mV for at least 30 s and stepping to +60 mV for 500 ms. All scale bars represent 200 pA and 20 ms.

**Channel Modulation by Kvβ Substrates**—That Kvβ is a functional AKR naturally leads to the question of whether the redox reaction on Kvβ affects channel function. To address this question, we co-expressed Kv1.4 with Kvβ2 and monitored channel function on inside-out patches before and after applying a Kvβ2 substrate.

Fig. 4A shows currents from a typical inside-out patch excised from an oocyte co-expressing Kv1.4 and Kvβ2. When 4-CY was perfused on the intracellular side of the membrane, channel inactivation rate decreased from 81 ± 1.9 s⁻¹ to 46 ± 1.4 s⁻¹ (Fig. 4, A and E, and Table 1; n = 31 patches from eight batches of oocytes), and both the peak current and the steady-state current levels increased significantly (Fig. 4A). 4-CB induced an almost identical response (data not shown). We have observed the substrate-induced change of channel current consistently on all the inside-out patches, and the change remained after the substrates were washed away.

When 4-CY was perfused to patches expressing only Kv1.4, the rate of channel inactivation changed slightly from 35 ± 2.3 to 34 ± 2.7 s⁻¹ (Fig. 4, B and E, and Table 1; n = 10 patches from four batches of oocytes), and the current level remained essentially unchanged, indicating that the substrates modulate channel function through Kvβ.

To find out if the change in channel function is due to the redox reaction on Kvβ, we tested 4-CY on Kv1.4 co-expressed with either the D85N or the K118M mutant Kvβ2. In the case of Kv1.4 paired with the D85N mutant, the inactivation rate constant changes from 61 ± 4.1 to 58 ± 4.0 s⁻¹ (Fig. 4, C and E, and Table 1; n = 9, four batches of oocytes). In the case of Kv1.4 paired with the K118M Kvβ2, it changes from 59 ± 2.4 to 57 ± 2.5 s⁻¹ (Fig. 4, D and E, and Table 1; n = 6, three batches of oocytes). In both cases, 4-CY only induced a small change in channel current level and altered the rate of inactivation slightly, indicating that 4-CY is less efficient in modulating channel inactivation when the mutant Kvβ is present. Combined, the electrophysiological studies indicate that the substrates modulate channel functions through Kvβ2 and most likely through inducing the redox reaction.

**Oxidation of the Bound NADPH Modulates Channel Function**—How does a Kvβ substrate modulate channel function? On an excised patch, Kvβ very likely contains an NADPH, and when a substrate is applied to the patch, the Kvβ-bound NADPH is oxidized. The NADPH-to-NADP⁺ conversion induces a change in channel inactivation. If this hypothesis is true, we can make the following two predictions and test them experimentally.

First, we reasoned that perfusing a patch with NADPH after channel modulation should reverse the 4-CY/4-CB effect because it would re-load Kvβ2 with the reduced cofactor. Indeed the 4-CY effect was slowly reversed when the patch was perfused with 0.2 mM of NADPH (Fig. 4F, left panel). Recovery of current was observed in all the patches (n = 8), and in two of them the current recovered to the level before 4-CY application. In sharp contrast, no change in channel current was observed when NADPH was perfused to patches (n = 5) expressing Kv1.4 only (Fig. 4F, right panel). These experiments suggest that Kvβ on an inside-out patch can complete an enzymatic cycle, when 4-CY was perfused to oxidize its bound cofactor and followed by fresh NADPH to re-prime the enzyme. Changes in channel function served as a readout.

Second, we found that when a patch expressing both Kv1.4 and Kvβ2 was exposed to a substrate, channel current increases gradually over time and reaches a steady state (Fig. 5A). This suggests that although we do not know how the NADPH-to-NADP⁺ conversion induces a change in channel current, we can nevertheless follow the time course of channel modulation by measuring the current level as a function of time after a substrate was applied. Since 4-CB reacts with the bound NADPH at a rate ~3-fold slower than 4-CY (Fig. 2C), we predict that channel modulation by 4-CB may also be slower if the cofactor oxidation is required for channel modulation. To test this prediction, we measured channel
Inactivation time constants for Kv1.4 channels paired with different Kvβ2s

Current decay was fit with an exponential function with two components, r1 and r2. The fraction of each component is f1 and f2, respectively. The Kv1.4 used in this study has a cysteine at position 13 mutated to a serine (see “Experimental Procedures”).

|        | m/s | r1            | r2            |
|--------|-----|---------------|---------------|
| Kv1.4  | 28.2 ± 1.8 | 0.54 ± 0.061  | 0.46 ± 0.061  |
| Kv1.4 (after 4-CY) | 29.6 ± 2.4 | 0.61 ± 0.054  | 0.39 ± 0.054  |
| Kv1.4 + Kvβ2 | 12.3 ± 0.28 | 0.81 ± 0.013  | 0.19 ± 0.012  |
| Kv1.4 + Kvβ2 (after 4-CY) | 21.7 ± 0.68 | 0.74 ± 0.021  | 0.26 ± 0.021  |
| Kv1.4 + Kvβ2 D8SN | 16.4 ± 1.1 | 0.80 ± 0.023  | 0.20 ± 0.023  |
| Kv1.4 + Kvβ2 D8SN (after 4-CY) | 17.3 ± 1.2 | 0.80 ± 0.031  | 0.20 ± 0.030  |
| Kv1.4 + Kvβ2 K118M | 17.0 ± 0.68 | 0.64 ± 0.054  | 0.36 ± 0.054  |
| Kv1.4 + Kvβ2 K118 M (after 4-CY) | 17.5 ± 0.76 | 0.71 ± 0.055  | 0.29 ± 0.055  |

Kvβ is a functional enzyme, it is now possible to test if other aspects of channel functions are directly coupled to the oxidation of the bound NADPH cofactor as well.

In a generic AKR, there are four highly catalytic residues (Fig. 1B). Of the four, the catalytic aspartate, tyrosine, and lysine are almost 100% conserved among all AKR families including Kvβ, while the histidine is conserved in some of the AKR families, and in Kvβ it is an asparagine. In a 3α-HSD when the histidine was mutated to a glutamate, it has little impact on the turnover rate constant but increased the Km by a factor of 10 (38). Similarly, mutating the histidine to an asparagine in an aldehyde reductase reduced the turnover rate constant by only a factor of 3 but substantially increased the Km value of the substrates (39). These results indicate that the histidine residue may play an important role in substrate recognition, and the corresponding asparagine residue in Kvβ could be evolved for the recognition of its specific native substrate.

The multiple enzymatic turnover rate constant of Kvβ2 is ~0.08 min⁻¹ and is almost the same for the two substrates. In comparison, 3α-HSD catalyzes a very similar substrate, 4-nitrobenzaldehyde, at a rate of 61.4 min⁻¹ (40), almost 760-fold faster than Kvβ2. On the other hand, the single turnover rate constant of Kvβ is at least 1.9 min⁻¹. Compared with a 3α-HSD, the native substrate 5α-dihydrotestosterone oxidizes the bound NADPH at a limiting rate of 26 min⁻¹ (41), which is ~14-fold faster than Kvβ. These results indicate that Kvβ is a slow enzyme mainly because it has a slow cofactor exchange rate.

Is Kvβ a redox sensor that detects change in cellular chemistry and modulates channel function or an enzyme whose catalytic activity is regulated by channel activities? Data presented here cannot distinguish between the two possibilities but seem to favor somewhat the redox sensor mechanism. The enzymatic turnover rate constant appears too slow if the function of Kvβ is to catalyze substrate turnover but not so if it is to sense cellular redox changes, which would require only the hydride transfer step and occurs over the time scale consistent with our experiments. Furthermore, earlier structural studies have shown that the bound NADPH was oxidized over a period of 2 weeks during the crystallization process, but nevertheless the NADP⁺ still was present in the structure of Kvβ2 (10). The tight association is partly due to a flexible loop stretching over the NADPH binding site and restricting the hydride transfer step and occurs over the time scale consistent with our experiments. Furthermore, earlier structural studies have shown that the bound NADPH was oxidized over a period of 2 weeks during the crystallization process, but nevertheless the NADP⁺ still was present in the structure of Kvβ2 (10). The tight association is partly due to a flexible loop stretching over the NADPH binding site and restricting the hydride transfer step and occurs over the time scale consistent with our experiments. Furthermore, earlier structural studies have shown that the bound NADPH was oxidized over a period of 2 weeks during the crystallization process, but nevertheless the NADP⁺ still was present in the structure of Kvβ2 (10).
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require understanding the detailed mechanisms of the functional coupling and discovering the physiological substrate(s) of Kvβ.

The functional interactions between the two proteins demonstrate a simple way of coupling intracellular redox states to the excitability of a cell. The results presented here bring us one step forward to understanding the physiological functions of Kvβ.

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