Mechanisms of Barbiturate Inhibition of Acetylcholine Receptor Channels

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ABSTRACT We used patch clamp techniques to study the inhibitory effects of pentobarbital and barbital on nicotinic acetylcholine receptor channels from BC3H-1 cells. Single channel recording from outside-out patches reveals that both drugs cause acetylcholine-activated channel events to occur in bursts. The mean duration of gaps within bursts is 2 ms for 0.1 mM pentobarbital and 0.05 ms for 1 mM barbital. In addition, 1 mM barbital reduces the apparent single channel current by 15%. Both barbiturates decrease the duration of openings within a burst but have only a small effect on the burst duration. Macroscopic currents were activated by rapid perfusion of 300 μM acetylcholine to outside-out patches. The concentration dependence of peak current inhibition was fit with a Hill function; for pentobarbital, $K_i = 32 \mu M, n = 1.09$; for barbital, $K_i = 1900 \mu M, n = 1.24$. Inhibition is voltage independent. The kinetics of inhibition by pentobarbital are at least 30 times faster than inhibition by barbital (3 ms vs. <0.1 ms at the $K_i$). Pentobarbital binds ~10-fold more tightly to open channels than to closed channels; we could not determine whether the binding of barbital is state dependent. Experiments performed with both barbiturates reveal that they do not compete for a single binding site on the acetylcholine receptor channel protein, but the binding of one barbiturate destabilizes the binding of the other. These results support a kinetic model in which barbiturates bind to both open and closed states of the AChR and block the flow of ions through the channel. An additional, lower-affinity binding site for pentobarbital may explain the effects seen at >100 μM pentobarbital.

KEY WORDS: pentobarbital • barbital • anesthetic • patch clamp • ion channels

INTRODUCTION Barbiturates have several therapeutic effects on humans including light sleep (at low doses), deep coma (at high doses), amnesia, muscle relaxation, protection against cerebral ischemia, and reversal of seizures (Fragen, 1994). Thus, these drugs probably have multiple effects on the central and peripheral nervous systems. Ion channels are among the possible targets of barbiturates. Barbiturates are known to affect many types of ion channels including GABA<sub>R</sub> receptor channels (Tanelian et al., 1993), some (ffrench-Mullen et al., 1993) but not all (Hall et al., 1994) calcium channels, sodium channels (Frenkel et al., 1990; Barann et al., 1993), glutamate receptor channels (Marszalec and Narahashi, 1993), 5-HT<sub>3</sub> receptor channels (Barann et al., 1993), and muscle-type nicotinic acetylcholine receptor (AChR)<sup>1</sup> channels (deArmendi et al., 1993).

<sup>1</sup>Abbreviations used in this paper: ACh, acetylcholine; AChR, ACh receptor; Barb, barbital; PB, pentobarbital.

Inhibition of AChR by barbiturates has been studied with electrophysiological (Lee-Son et al., 1975; Gage and McKinnon, 1985; Jacobson et al., 1991; Yost and Dodson, 1993), flux (Firestone et al., 1986; Roth et al., 1989; deArmendi et al., 1993), and binding (Dodson et al., 1990) techniques. Several effects have been observed (Firestone et al., 1986; 1994), but the dominant effect is a direct inhibitory action of the drug on the open state of the channel. The potency of barbiturates for inhibiting the channel is related to but not completely determined by lipid solubility (deArmendi et al., 1993). A study of the single channel kinetics in the presence of pentobarbital (PB) allowed Gage and McKinnon (1985) to dismiss a simple, sequential open channel blocking mechanism for PB action. They suggested that the mechanism might be allosteric in that the binding of one molecule of PB to the open channel protein induces a conformational change to a new closed state of the channel. In this scenario, the binding of PB is not concomitant with inhibition.

Here, we use several patch clamp recording protocols to study the effects of PB and barbital (Barb) on nicotinic AChRs in outside-out patches from BC3H-1 cells. We examine single channel kinetics, the equilibrium and kinetic properties of macroscopic currents,
and interactions between PB and Barb. We conclude that inhibition of the AChR by barbiturates is temporally coincident with binding of the drug to a site on the channel protein. This can be described by a model in which barbiturates bind to both the open and closed states of the channel. PB shows a strong preference for binding to the open state. The binding sites for PB and Barb do not coincide but are probably close to each other. There is evidence for an additional, lower affinity binding site for PB.

**Materials and Methods**

BC3H-1 cells that express the α3β4α6-type nicotinic AChR were cultured as described previously (Sine and Steinbach, 1984). To prepare cells for patch clamp recording, the culture medium was replaced with an “extracellular” solution (ECS) containing (in mM): NaCl (150), KCl (5), CaCl₂ (1.8), MgCl₂ (1.0), and HEPES (10), pH 7.3. Patch pipettes were filled with a solution consisting of (in mM) KCl (140), EGTA (5), MgCl₂ (5), and HEPES (10), pH 7.3, and had resistances of 3–6 MΩ. ECS. The 20 mM Barb solution had a pH of 8.3; this was titrated to 20 mM Barb or 1 mM PB (Sigma Chemical Co., St. Louis, MO) in ECS. The 20 mM Barb solution had a pH of 8.3; this was titrated to pH 7.3, and had resistances of 3–6 MΩ. An outside-out patch (Hamill et al., 1981) with a seal resistance of 10 GΩ or greater was obtained from a cell and moved into position at the outflow of a perfusion system. The perfusion system consisted of solution reservoirs, manual switching valves, and a V-shaped piece of plastic tubing inserted into the culture dish (Liu and Dilger, 1991). For single channel current measurements, the manual valves were used to switch from drug-free to drug-containing solutions containing 0.2 mM ACh. For macroscopic current measurements, the perfusion system also contained a solenoid-driven pinch valve. One arm of the “V” contained ECS without agonist (normal solution); the other arm contained ECS with 60 individual current traces. Mean currents were fit to a 1- or 2-exponential function.

\[ I(t) = I_p + I_{1exp} \left( \frac{t}{\tau_1} \right) + I_{2exp} \left( \frac{t}{\tau_2} \right). \]  

The time constant of the 1-exponential fit and the slower time constant of the 2-exponential fit, \( \tau_1 \), measures the current decay due to desensitization (Dilger and Liu, 1992). In the presence of PB, the current contained an additional fast component, \( \tau_2 \). This represents the time course of inhibition by PB (see results). Fractional inhibition of the peak mean current, \( I_{\text{p,mean}} \), the maximum inward current obtained after perfusing ACh, was calculated as the ratio of the current in the presence of drug, \( I_{\text{p,mean}} \), to the current in the absence of drug, \( I_p \). For PB experiments, \( I_{\text{p,mean}} \) was obtained from the amplitude of the slow component of the decay, \( I_{\text{p,mean}} = I_p \) (see results).

Single channel recordings were made while the patch was exposed to ECS + 0.2 μM ACh at a patch potential of -100 mV. Data was digitized in 5-s segments at a rate of 50 μs per point. 3–10 data segments were collected (enough to obtain 200–1,000 single channel events, depending on the channel activity in the patch). Data collection was repeated with ECS + ACh + barbiturate and then again with ECS + ACh (recovery). Data were accepted if, after analysis, we found that the channel kinetics during recovery were within 20% of the values obtained during the initial data collection segments.

Single channel analysis consisted of identifying opening and closing transitions, obtaining the distribution of open, closed, and burst durations, and fitting the distributions (expressed as the number of events vs. log-binned duration, 10 bins per decade) to 1- and 2-exponential probability distribution functions by finding the maximum log-likelihood using a simplex algorithm. The single-exponential fit was considered adequate when the fractional amplitude of one of the components of the 2-exponential fit was <0.1. The definition of bursts was based on the distribution of short (gap) and long closures (Colquhoun and Sakmann, 1985). Mean gap and open durations and the number of openings per burst were corrected for undetected events using equations derived for a two-state mechanism (Colquhoun and Hawkes, 1995a). This approximation is probably adequate because of the large time separation between brief and long closed durations and our observation that the open time histogram in the presence of barbiturate usually has only one component. Mean single channel amplitudes were calculated by taking the average of the amplitudes of those openings lasting >0.25 ms; these are not attenuated by the 3 kHz low-pass filter.
**RESULTS**

The Effects of Barbital and Pentobarbital on AChR Single Channel Currents

The effects of Barb and PB on single AChR channels are illustrated in Fig. 1. Inward single channel currents at $-100 \text{ mV}$ were activated by a low concentration of ACh (0.2 $\mu$M). Under control conditions (Fig. 1, left), channel activity consists of $-4 \text{ pA}$ openings lasting an average of 4 ms and separated by tens of milliseconds. Occasionally, a brief closing transition is seen. Both 100 $\mu$M PB and 500 $\mu$M Barb (Fig. 1, right) induce a bursting pattern of channel activity. The closures within a burst are much longer for PB than for Barb. Barb-induced closures are so brief that the single channel amplitude appears to be attenuated by $\sim 8\%$. The attenuation is more pronounced at 1,000 $\mu$M Barb (see Fig. 5 D).

The closed duration histograms constructed from single channel data have two components (Fig. 2). The dominant component of the control histogram occurs near 60 ms and corresponds to the time between activation of different channels in the patch. For 0.2 $\mu$M ACh, a long closed time of 60 ms indicates that there are $\sim 150$ active channels in the patch. The small, brief component of the control histograms occurring at $<100 \mu$s most likely corresponds to the closing of a channel followed by the rapid reopening of the same channel. The bursting activity of the barbiturates is represented by a large number of gaps: the brief component of the closed time histogram. PB induces gaps near 2 ms; Barb induces gaps near 70 $\mu$s. Neither barbiturate has a significant effect on the long closed component.\(^3\)

The barbiturates decrease the open duration of single AChR channels but have little effect on the burst duration (Fig. 3). Under control conditions, there are

\(^2\)This estimate comes from comparison of single channel burst frequency with peak macroscopic currents measured on the same patch (Liu, Y., and J.P. Dilger, unpublished data).

\(^3\)For the data shown in Fig. 2 with 100 $\mu$M PB, the long closed component increased to 140 ms. However, subsequent return to control conditions showed the long closed time to be 140 ms. We assume that, in this patch, there was a rundown in channel activity between the first control and 100 $\mu$M PB runs. In patches that did not exhibit any rundown, 100 $\mu$M PB did not affect the long closed time. We did observe that 250 and 500 $\mu$M PB tended to decrease the long closed interval.

![Figure 1](image1.png)

**Figure 1.** Single AChR channels in control and in the presence of 100 $\mu$M PB or 500 $\mu$M Barb. Channels activated by 0.2 $\mu$M ACh, $-100 \text{ mV}$, two separate outside-out patches. Both barbiturates induce a bursting behavior of the channels but the closures within the bursts are longer for PB than for Barb. Patches L82 and L105.

![Figure 2](image2.png)

**Figure 2.** Closed duration histograms compiled from the same two experiments exemplified in Fig. 1. The solid lines are fits to the two-exponential probability distribution function. The long component represents the closed time between openings of different channels in the patch. The brief component represents gaps within a burst of activity of a single channel. Gaps are rarely seen in control recordings but are frequently seen in the presence of a barbiturate. The vertical calibration bar represents 10 events in control (linear scale), see below for vertical scaling of barbiturate histograms. 0.2 $\mu$M ACh, $-100 \text{ mV}$. (A) Control: 236 events, gap duration $= 66 \mu$s (fraction $= 0.12$); long duration 63 ms. 100 $\mu$M PB: 251 events, gap duration $= 1.8 \mu$s (0.48); long duration $= 140 \text{ ms}$. Calibration bar $= 6$ events. (B) Control: 509 events: gap duration $= 66 \mu$s (0.64); long duration $= 66 \text{ ms}$, calibration bar $= 50$ events.
two components in the open duration histogram; a brief one at 200–500 μs comprising 20–50% of the events and a long one at 4–5 ms. Because there are very few brief closures within a burst, the control burst duration histogram is very similar to the control open duration histogram. In the presence of either 100 μM PB or 500 μM Barb, the open duration histogram collapses to a single component with a time constant of 1.2–1.4 ms. In contrast, there is very little difference between the control and barbiturate burst duration histograms. The vertical calibration bar represents 10 events in control (linear scale), see below for vertical scaling of barbiturate histograms. 0.2 μM ACh, −100 mV. (A) Open durations. Control: 226 events, $\tau_{\text{fast}} = 0.48$ ms (.35), $\tau_{\text{slow}} = 4.9$ ms. 100 μM PB: 251 events, $\tau = 1.3$ ms, calibration bar = 14 events. (B) Open durations. Control: 304 events, $\tau_{\text{fast}} = 0.26$ ms (.34), $\tau_{\text{slow}} = 4.1$ ms. 500 μM Barb: 723 events, $\tau = 1.2$ ms, calibration bar = 25 events. (C) Burst durations. Control: 212 events, 1.07 openings per burst, $\tau_{\text{fast}} = 0.96$ ms (.46), $\tau_{\text{slow}} = 6.3$ ms. 100 μM PB: 136 events, 1.85 openings per burst, $\tau_{\text{fast}} = 0.21$ ms (.25), $\tau_{\text{slow}} = 4.7$ ms, calibration bar = 7 events. (D) Burst durations. Control: 294 events, 1.03 openings per burst, $\tau_{\text{fast}} = 0.23$ ms (.33), $\tau_{\text{slow}} = 4.2$ ms. 500 μM Barb: 380 events, 1.90 openings per burst, $\tau_{\text{fast}} = 0.16$ ms (.28), $\tau_{\text{slow}} = 3.3$ ms, calibration bar = 13 events.
used. To verify that the correction procedure for accounting for missed events provides good estimates of the mean open and gap durations, we studied six patches using a cutoff frequency of 6 kHz, a sampling time of 25 μs, and an applied voltage of −125 mV. Under these conditions, there was very little variation of the current amplitude with Barb concentration; the amplitude was 4% lower with 1 mM Barb than with control. The gap duration remained 50 μs. At the 6 kHz resolution, a greater number of gaps are resolved, but after correcting for undetected events, the values of \( N_{\text{open/burst}} \) and \( \tau_{\text{open}} \) are no different than at the 3 kHz resolution. The Barb concentration dependence of the burst duration was the same for both the 3 kHz and 6 kHz data.

If we assume that the Barb-induced bursts are composed of brief openings to the fully opened state and brief closures to the fully closed state, we can use a beta function analysis of the amplitude histogram to estimate the open and closed time within bursts (Yellen, 1984). To do this, we applied a 1-kHz Gaussian digital filter to the single channel data and constructed an amplitude distribution from segments containing single bursts. This distribution is then fit to a beta function containing two parameters: the average open and closed times within bursts. For four patches with 1 mM Barb, the average open time was 150 ± 70 μs, and the average closed time was 46 ± 17 μs. This estimate of the open time is shorter than the one obtained by analyzing single channel data with 1 mM Barb (540 ± 110 μs, Fig. 5 A), but the estimate of the closed time is similar to the average gap duration from single channel analysis. This suggests that even after correcting the single channel data for unresolved events, we may overestimate the open time and underestimate the number of openings per burst.

The single channel data suggest that both PB and Barb act, at least qualitatively, as blockers of the AChR channel. In this interpretation, the two barbiturates differ in the duration of blocking events: the less potent drug, Barb, blocks for <0.1 ms, and the more potent drug, PB, blocks for ~2 ms. In the discussion, we make a quantitative test of models in which barbiturates block both open and closed AChR channels. Be-

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**Figure 4.** The PB concentration dependence of channel kinetics and amplitudes. (A) Open and burst durations (longer components are used when two components are present). (B) The number of openings per burst. (C) The duration of gaps between bursts. (D) The single channel current amplitude. The solid lines represent the predictions of scheme i with the kinetic parameters given in Table I. The dashed lines represent predictions of scheme iii for burst duration, gap duration, and openings per burst. 0.2 μM ACh, −100 mV.
fore doing so, we present data from macroscopic current experiments that provide additional information about the action of barbiturates on the channel.

The Effects of Barbital and Pentobarbital on Macroscopic AChR Currents

Both Barb and PB inhibit the macroscopic currents evoked by rapid perfusion of ACh. Fig. 6 presents examples of currents activated by 300 μM ACh in control and in the continuous presence of 50 and 400 μM PB (Fig. 6 A) or 2 and 20 mM Barb (Fig. 6 B). In the control traces, the current reaches a peak within <1 ms and then decays with a time constant of 50–60 ms due to desensitization. With 2 or 20 mM Barb, the peak currents are reduced to 60 or 5% of control and desensitization occurs with the same time course as in control. It appears that Barb interacts with the channels either before they are opened by ACh or very quickly after ACh is perfused. With 50 or 400 μM PB, an initial fast decay precedes desensitization. This suggests that, unlike Barb, PB is not very effective at interacting with closed channels. We explore this in more detail below. To determine the degree of inhibition of open channels by PB, we fit the data to a 2-exponential decay and extrapolate the slow component (desensitization) to $t = 0$ (Fig. 6 B, dotted lines). The extrapolated peak currents are reduced to 35% (50 μM) or 4% (400 μM PB) of control.

The results from experiments on nine patches with Barb (relative peak currents) and six patches with PB (relative extrapolated peak currents) are summarized in Fig. 7. These data were fit to the Hill equation (Fig. 7, solid lines):

$$\frac{I_p}{I_p^0} = \frac{K_i^n}{K_i^n + [B]^n},$$

where $[B]$ is the drug concentration, $K_i$ is the drug concentration needed for 50% inhibition (IC$_{50}$), and $n$ is the Hill coefficient. For Barb: $K_i = 1.9 \pm 0.2$ mM, $n = 1.24 \pm 0.07$; for PB: $K_i = 32 \pm 2$ μM, $n = 1.09 \pm 0.06$. Thus, PB is 60 times more potent than Barb at inhibiting the AChR. Because the Hill slopes are close to unity, it is possible that only one barbiturate molecule is involved in the inhibition of a channel.

The fast decay that occurs in the macroscopic currents with PB provides information about the rate of equilibration of PB with the channel. The decay is
faster with higher concentrations of PB. In Fig. 6 A, the time constants are 3.2 and 1.9 ms for 50 and 400 μM PB, respectively. Fig. 8 shows that the fast decay time decreases monotonically with the concentration of PB.

The inhibitory effect of PB is not voltage dependent. This is illustrated in Fig. 9 in which ACh-activated currents at three different voltages, −100, −50, and +50 mV, are compared for a single patch. The traces are scaled so that the controls have the same peak current level. Neither the time constant of the fast current decay, nor the extrapolated level of the residual current is affected by voltage. There was no difference in the effect of Barb on macroscopic currents over this voltage range either (not shown).

The macroscopic current data used for Figs. 6–9 were obtained with equilibrium drug concentrations; that is, the barbiturate was present in both the normal and test perfusion solutions. Information about the kinetics of current inhibition by the barbiturates can be obtained from experiments in which the drug is applied simulta-
neously with ACh. Fig. 10 shows that, for 100 μM PB, the current decay under equilibrium conditions and the current decay after rapid addition of the drug (onset) are identical. This supports the idea that PB has very little interaction with channels when they are closed; prior exposure to PB does not change the degree of inhibition either at the peak or during the decay of the current response.

Rapid addition of Barb also produces a current decay, but on a much faster time scale (Fig. 11). For this experiment, the time resolution was increased by perfusing with 10 mM ACh (this saturates the ACh binding sites within microseconds so that the 20–80% risetime of the current, 40 μs, is determined mainly by the channel opening rate [Liu and Dilger, 1991]), filtering at 15 kHz, sampling at 5 μs per point, and using +50 mV to avoid channel block by ACh. Two pieces of qualitative information about the effects of 5 mM Barb can be extracted from Fig. 11. The equilibrium trace shows that, in contrast to PB, the inhibitory effect of Barb is present at all times after perfusion with ACh. We conclude that Barb either interacts with closed channels to the same degree as it interacts with open channels or, it does not interact with closed channels but equilibrates with open channels extremely quickly, on the order of tens of microseconds or faster. The second observation, that the onset current trace exhibits a 60-μs decay, probably reflects both the binding kinetics of Barb and the time course of the Barb concentration jump. Similarly, the kinetics of recovery from block by Barb show a relaxation from the equilibration level of inhibition to control with a time course of 50 μs (not shown). The time resolution of these experiments is not sufficient to determine if this represents the kinetics of Barb dissociation from its inhibitory site or simply the diffusion of Barb away from the patch.

**Interactions between Barbital and Pentobarbital**

To determine whether PB and Barb inhibit the AChR channel by binding to a single site on the AChR protein, we performed experiments with both barbiturates. Fig. 12 is an example with 5 mM Barb and 100 μM PB.
In the left panel, 100 μM PB decreases the extrapolated peak current to 22% of control. In the right panel, 5 mM Barb decreases the current to 34% of control (note different current scale). When both barbiturates are present, the current decreases to 40% of the 5 mM Barb current (14% of control). Thus, PB is less effective when applied in combination with Barb than when applied by itself. If the two drugs acted independently, the current would have been 22 × 34% or 7.5% of the control. If the two drugs compete for the same binding site, the predicted current is 15% of control (see discussion). The time constant of the fast decay component is 1.6 ms for PB alone and 1.8 ms for both barbiturates in combination.

Inhibition curves for PB alone and PB + 5 mM Barb are shown in Fig. 13. The latter data are normalized to the relative current observed in the presence of 5 mM Barb alone, 0.30 ± 0.02 (n = 25). Fits of the data to the Hill equation (Eq. 2), give $K_i = 28 ± 2 \mu M$ for PB alone and $K_i = 53 ± 3 \mu M$ for PB + 5 mM Barb. For both data sets, the Hill coefficient was close to unity: 1.09 ± 0.07 and 0.96 ± 0.07, respectively. Thus, there is a considerable decrease in the effectiveness of PB when Barb is present. However, this decrease is not as great as would be expected if PB and Barb were competing for a single binding site (Fig. 13, dotted line; see discussion). In the presence of 5 mM Barb, the time constant of the fast decay seen with PB is decreased at 25 and 50 μM [PB] but is unchanged at [PB] ≈ 100 μM (Fig. 14).
**Figure 14.** The PB concentration of $\tau_{\text{onset}}$ obtained from the fast decay component of macroscopic currents in the absence (open symbols) and presence (closed symbols) of 5 mM Barb. The solid lines are fits of the data to Eq. 8 with $f = 4.8 \times 10^6 / \text{M/s}$ and $b = 200 / \text{s}$ (PB alone) and $f = 3.5 \times 10^6 / \text{M/s}$ and $b = 340 / \text{s}$ (PB + 5 mM Barb). In the presence of Barb, PB binds less quickly and dissociates more quickly, 300 $\mu$M ACh, -50 mV.

### DISCUSSION

**Effects Seen with One Barbiturate**

The bursting effect of the barbiturates on single AChR channels suggests a model in which drug molecules bind to the AChR and block the flow of ions through the channel. This bursting cannot be explained by a model in which drug molecules bind to and block only the open state of the AChR (purely open channel block) because the expected increase in burst duration (Neher, 1983) is not seen (Figs. 4A and 5A). We will test the adequacy of the model shown as **Scheme 1** (Murrell et al., 1991; Dilger et al., 1992) in which barbiturate molecules ($B$) can bind to both the open ($O$) and closed ($C$) conformations of the channel.

$$
\begin{align*}
C & \xleftarrow{\beta} \xrightarrow{\alpha} O \\
& \xleftarrow{b'} f[B] \xrightarrow{b} f[B] \\
& \xleftarrow{\beta'} \xrightarrow{\alpha'} CB \xrightarrow{OB} \text{(**Scheme 1**)}
\end{align*}
$$

In **Scheme 1**, the various closed states of the AChR (unliganded, singly liganded, and doubly liganded) are represented by a single state. The effective opening rate, $\beta$, depends on agonist binding, agonist concentration, and the channel opening rate. The channel closing rate is $\alpha$. A barbiturate molecule may bind to either the closed state (to form CB) or open state (to form OB); the association ($f$ and $f'$) and dissociation ($b$ and $b'$) rates may depend on the channel conformation. The gating transition rates between drug-bound open and closed states ($\alpha'$ and $\beta'$) may differ from the normal gating transition rates. In **Scheme 1**, a single barbiturate molecule is sufficient to inhibit one channel; this is supported by the concentration-inhibition curves for PB and Barb (Fig. 7) that have Hill coefficients close to unity.

**Scheme 1** makes quantitative predictions about the drug concentration dependence of $\tau_{\text{open}}$, $\tau_{\text{burst}}$, $\tau_{\text{gap}}$, and $N_{\text{open/burst}}$ (Dilger et al., 1992).

$$
\begin{align*}
\tau_{\text{open}} &= \frac{1}{\alpha + f[B]}, \\
\tau_{\text{burst}} &= \frac{1 + f[B]b}{\alpha + f[B]b' / (b + \alpha')}, \\
\tau_{\text{gap}} &= \frac{1}{b + \alpha'}, \\
N_{\text{open/burst}} &= \frac{\alpha + f[B]}{\alpha + f[B]b' / (b + \alpha')}.
\end{align*}
$$

The relative peak current induced by rapid perfusion of saturating concentrations of ACh can also be calculated from **Scheme 1**.

$$
\frac{I_r}{I_p} = \left( 1 + \frac{f[B]}{b} \right)^{-1}.
$$

**Scheme 1** predicts that the time constant of the current decay induced by a jump in drug concentration is:

$$
\tau_{\text{onset}} = \frac{1}{f[B] + b}.
$$

Note that, in **Scheme 1**, the time constant of the macroscopic current decay is dependent on the same parameters as the kinetics of channel flickering. Hence, the decay is the multi-channel correlate of single channel flickering. Neither $\alpha$ nor $\alpha'$ appear in the macroscopic current expressions (Eqs. 7 and 8) because, with saturating concentrations of ACh, dissociation of one molecule of ACh is quickly followed by binding of another. Under these conditions, the concept of burst loses its meaning.

The $K_i$ values (from Fig. 7) for PB (32 $\mu$M) and Barb (1.9 mM) determine the equilibrium between open and open-blocked channels ($b/f$) in **Scheme 1**. These values are about twofold greater than those reported for PB and Barb inhibition of flux in *Torpedo* AChR (deArmendi et al., 1993). For PB, the association rates...
are given by fitting the concentration dependence of the single channel open time (Eq. 3; Fig. 4 A, solid line); \( f = 6.5 \times 10^6/M/s \). The dissociation constant can be calculated from \( f \times b/f \); \( b = 210/s \). Very similar values for the association and dissociation constants for PB are obtained by fitting the concentration dependence of the fast decay time constant (Eq. 8, Fig. 8); \( f = 4.0 \pm 0.6 \times 10^6/M/s \) and \( b = 210 \pm 20/s \).

For Barb, analysis of the amplitude distribution suggests the open duration is more severely affected by the limited time resolution of the recording system than are the gap durations. Assuming for the moment that \( b >> \alpha' \), the observed gap duration gives \( b = 2 \times 10^4/s \) (Eq. 5, Fig. 5 C). Combining this with the equilibrium dissociation constant gives \( f = 1 \times 10^7/M/s \). This is faster than the value obtained by fitting the Barb concentration dependence of the open duration (\( f = 4 \times 10^6/M/s \)) but within the range obtained from analysis of the amplitude distribution (6–12 \( \times 10^6/M/s \)).

Estimates for the remaining undetermined parameter in scheme 1, \( \alpha' \), can be obtained by fitting the concentration dependence of either the burst duration or the number of openings per burst. However, the burst duration may be the better measurement to fit because unresolved events will affect the number of openings per burst more than the burst duration. For PB, the burst duration is very sensitive to the value of \( \alpha' \); only values in the range 150–220/s provide a good description of the data at \( \leq 100 \mu M \) PB. With \( \alpha' = 200/s \), the predicted number of openings per burst do not differ very much from the observed values (Fig. 4 B, solid line). For Barb, \( \alpha' \) is not as well defined; values in the range 100–800/s all predict a fairly flat concentration dependence of the burst duration. With all of these values of \( \alpha' \), the predicted number of openings per burst is much higher than the observed values at 1,000 \( \mu M \) Barb; \( N_{open/burst} = 32 \) with \( \alpha' = 100/s \) and 16 with \( \alpha' = 800/s \). This range of values for \( \alpha' \) satisfies the assumption that \( b >> \alpha' \), validating the estimate of \( b \) from the gap duration. The predictions of scheme 1 are shown with solid lines in Figs. 4 and 5 using the best fitting values (or intermediate values when there is a range of acceptable values) of \( f, b, \) and \( \alpha' \) (Table I).

In their single channel study of PB on ACh receptors in denervated mouse muscle, Gage and McKinnon found quantitatively similar results on the open, gap, and burst durations. From the concentration dependence of the open duration, they calculated \( f = 3.4 \times 10^6/M/s \) (16°C). They found a fivefold increase in the gap duration over the range of 10–500 \( \mu M \) PB from 1 to 5 ms. They considered this latter result as definite evidence against a sequential open channel blocking mechanism (scheme 1 without the CB state) but did not explore any additional models.

A state dependence for barbiturate binding to Torpedo AChRs was observed by deArmendi et al. (1993). They found that the open state is preferred over the closed state by a factor of 4.7 (PB) and 3.2 (Barb). These values were determined by comparing the concentration of barbiturate needed to inhibit flux with the concentration needed to displace \( [\text{H}] \)amobarbitral bound to the resting receptor (Dodson et al., 1990). The \( \sim 100 \mu s \) time resolution of our patch clamp experiments limits our ability to quantify the degree of barbiturate binding to the closed state. Fig. 10 indicates that there is no more than a 10% block of the closed channel with 100 \( \mu M \) PB. This implies a binding affinity to the closed state on the order of 1 mM and an open/closed state preference of about 30-fold for PB in AChRs from BC3H-1 cells. We cannot determine the state preference of Barb for our experiments (Fig. 11).

### Interactions between Barbital and Pentobarbital

The experiments illustrated in Figs. 12–14 address the question of whether PB and Barb compete for a single binding site on the AChR channel. If binding of the two drugs were absolutely competitive, the inhibition curve for PB in the presence of Barb, would be described by Eq. 9.

\[
\frac{I_p}{I_p^*} = \frac{1}{1 + \frac{[\text{PB}]}{K_{PB}} + \frac{[\text{Barb}]}{K_{Barb}}}.
\]

With 5 mM Barb, Eq. 9 predicts a 3.3-fold shift ([Barb]/\( K_{\text{Barb}} \)) of the PB inhibition curve to a half maximum effect at 94 \( \mu M \) PB (Fig. 13, dashed line). The observed shift of the half maximum concentration is only 1.9-fold. Thus, the binding of PB does not exclude the binding of Barb. The binding of the two drugs is not independent either. The presence of Barb decreases the binding affinity of PB. This is also apparent from measurements of onset kinetics (Fig. 14). In the presence of Barb, PB exhibits faster kinetics. Fits of the data to Eq. 8 indicate that, in the presence of 5 mM Barb, the association rate of PB is decreased (from 4.8 \( \pm 0.6 \times 10^6/M/s \) to 3.5 \( \pm 0.8 \times 10^6/M/s \)), and the dissociation rate of PB is increased (from 200 \( \pm 25/s \) to \( b = 340 \pm 70/s \)). One interpretation is that the binding sites for PB and Barb are the same but when both drugs bind, they have to move to nearby, less stable positions. Alter-

### Table I

| Scheme | Pentobarbital | Barbital |
|--------|---------------|----------|
| \( f (M^{-1}s^{-1}) \) | \( 6.5 \times 10^6 \) | \( 1.0 \times 10^7 \) |
| \( b (s^{-1}) \) | \( 2.5 \times 10^6 \) | \( 2.0 \times 10^4 \) |
| \( \alpha' (s^{-1}) \) | \( 2.0 \times 10^6 \) | \( 4.0 \times 10^8 \) |
| \( b/f (M^{-1}) \) | \( 3.8 \times 10^{-3} \) | \( 2.0 \times 10^{-3} \) |

The rate constants for scheme I used in fitting the single channel data (solid lines in Figs. 4 and 5). 0.2 \( \mu M \) ACh, –100 mV.
natively, there could be two distinct binding sites for the drugs and these sites interact allosterically. Our data cannot distinguish between these two possibilities.

**Allosteric Model**

An alternative interpretation of the bursting behavior induced by barbiturates is to consider bursting to arise from the control burst activity at rates modified by barbiturates. **Scheme II** is a model that is often used to describe the normal kinetics of AChR single channels (Auerbach, 1993).

\[
R \xrightarrow{\alpha} AR \xrightarrow{k_{-2}} A_2R \xrightarrow{\beta} A_2R^* \tag{Scheme II}
\]

In this scheme, R represents the receptor and A represents ACh. Channel activation results from the binding to two molecules of ACh followed by a conformational change from the doubly-ligated closed state (A2R) to the open state (A2R*). At low concentrations of ACh, a burst consists of one or more transitions between A2R and A2R* terminated by the dissociation of the agonist at a rate k_{-2}. The open time is given by 1/\alpha, the gap duration by 1/(\beta + k_{-2}) and the number of gaps per burst by \beta/k_{-2}. Under control conditions, \alpha = 0.5/\text{ms}, \beta = k_{-2} = 30/\text{ms} (Auerbach, 1993), so that the average burst consists of two 2-ms openings and one 20-\mu s gap (with the time resolution of our experiments, very few of these gaps are detected). Assume that the binding of barbiturates (with microsecond kinetics) modifies these rates to produce the observed burst kinetics. We have calculated the rates at each concentration of barbiturate. For both PB and Barb, \alpha increases as a function of concentration and is on the order of 1/\text{ms} for 100 \mu M PB and 250 \mu M Barb. 100 \mu M PB decreases both \beta and k_{-2} by a factor of 100. The effects of 250 \mu M Barb are more moderate; \beta decreases by a factor of 2 and k_{-2} decreases by a factor of 5. The difficulty with scheme II, however, is that it cannot account for the fast decay seen in macroscopic currents with rapid perfusion of 300 \mu M ACh in the presence of PB (Fig. 10). Moreover, scheme II predicts that the onset of macroscopic currents would have an onset time (at high concentrations of ACh) of 1/(\alpha + \beta), which is predicted to be 0.8 ms at 100 \mu M PB. Experimentally, we do not see any decrease in onset time (Fig. 10). We conclude that an allosteric model such as scheme II is not viable explanation for the effects of barbiturates.

**Extension of the Blocking Model**

In scheme I, the single channel gap duration (Eq. 5) is inversely proportional to the sum of b and \alpha' and is independent of the barbiturate concentration. For PB, the predicted gap duration is 2.2 ms (Fig. 4 C, solid line). The observed gap durations vary from 1.1 to 3.4 ms. The measured burst durations at high concentrations of PB also differ from the predictions of the model. One could argue that these deviations from the predicted values are unimportant because they occur at concentrations greater than three-times the K_i (after all, even the archetypal AChR open channel blocker, QX-222, shows deviations from predictions at high concentrations [Neher, 1983]). However, we wanted to determine whether we could use this information to gain further insights into the mechanism of action of the barbiturates. Several pieces of evidence suggest that the observed deviations from scheme I may be due to the binding of a second molecule of PB: (a) an increase in gap duration with [PB] is expected if the second molecule binds with low affinity and postpones the re-opening (unblocking) of the channel, (b) the interactions seen when both PB and Barb are present suggest that two barbiturate molecules may bind simultaneously, and (c) when the macroscopic current inhibition data (Fig. 7) is fit to a two-site inhibition function (Eq. 10), the two binding affinities are K_1 = 34 \pm 3 \mu M and K_2 = 800 \pm 500 \mu M (the dashed line in Fig. 7 is the prediction for two binding sites with affinities of 38 and 460 \mu M).

\[
\frac{I_p'}{I_p} = \frac{K_1 K_2}{K_1 K_2 + K_1 [PB] + [PB]^2}. \tag{10}
\]

We then considered whether scheme III could be used to quantitatively predict the observed single channel gap and burst distributions. Scheme III contains one additional state with two barbiturate molecules bound. We used Mathematica (version 2.2; Wolfram Research, Inc., Champaign, IL) to numerically evaluate the relevant matrix operations (Colquhoun and Hawkes, 1995b) for this model.

\[
\text{C} \mathrel{\xleftrightarrow{\beta}} \mathrel{\xleftrightarrow{\alpha}} \text{O}
\]

\[
\text{b'} \mathrel{\xleftrightarrow{f'[B]}} \text{b} \mathrel{\xleftrightarrow{f[B]}} \text{OB} \mathrel{\xleftrightarrow{f_2[B]}} \text{OB}_2 \tag{Scheme III}
\]

Numerical evaluation of scheme III requires values for nine independent constants. The channel activation rate, \beta, which depends on ACh binding, channel isomerization and the number of active channels, was set to 0.02/s to correspond to a typical control inter-burst interval of 400 ms. The previously determined values of \alpha and f were used (Table I). The values of \alpha' and b were adjusted to account for the gap duration at low concentrations of PB; the best agreement was obtained with \alpha' = 400/s and b = 300/s. We assumed that the poor interaction of PB with closed channels results from a low association rate and a normal dissociation...
rate: $f' = f/10$, $b' = b$. Detailed balancing was used to evaluate $\beta'$. We also assumed that the binding of a second molecule of PB has a normal association rate and a fast dissociation rate: $f_2 = f$, $b_2 = 12 \times b$ (giving $b_2/f_2 = 460 \, \mu M$ which is near the lower limit of the range obtained from a two-site fit). The results of the numerical evaluation are shown with dashed lines in Figs. 4 A, B, and C.4 SCHEME III quantitatively predicts the PB concentration dependence of the gap and burst durations and the number of openings per burst (the prediction for open duration is not shown because it is identical to that of SCHEME 1). Similar results are obtained when different assumptions are used (second binding site having a slow association rate and a normal dissociation rate: $f_2 = f/12$, $b_2 = b$; binding to closed channel having a normal association rate and a fast dissociation rate: $f' = f$, $b' = 10^6 b$). As might be expected, as the affinity of the second binding site is decreased, higher concentrations of PB are needed to obtain comparable changes in the predicted gap and burst durations (e.g., the predictions for $b_2/f_2 = 920 \, \mu M$ at 1,000 $\mu M$ PB are similar to the predictions for $b_2/f_2 = 460 \, \mu M$ at 500 $\mu M$ PB). The predictions of SCHEME III for macroscopic currents and kinetics do not differ significantly from the predictions of SCHEME I.

Summary

The action of PB and Barb differs primarily in the dissociation rate; Barb dissociates 80 times faster than PB. This is a greater difference than would be expected if lipid solubility were the only factor that determines barbiturate potency; the octanol:buffer partition coefficients are 106 (PB) and 4.5 (Barb), giving a ratio of 23 (Firestone et al., 1986a). The same conclusion was reached, based on flux experiments with 14 barbiturates, by deArmendi et al. (1993). Our experiments suggest that the inhibitory binding site is not identical for PB and Barb. This is a plausible explanation for the poor correlation between potency and lipid solubility. Interestingly, the potency ratio for Barb and PB anesthesia in tadpoles is also large: 14.6 $mM/0.16 \, mM = 90$ (Lee-Son et al., 1975).

The kinetic experiments described here do not directly address the question of the location of the barbiturate binding site(s). The close temporal association between the duration of inhibitory events seen at the single channel level (the gap duration in Figs. 4 and 5) and the kinetics of macroscopic current inhibition after rapid perfusion of barbiturate (the onset time in Figs. 10 and 11) suggest that barbiturate binding and channel inhibition are inseparable. This favors a steric blocking mechanism over an allosteric effect. This has also been observed with other anesthetics acting on the AChR channel (Dilger et al. 1994). Allosteric mechanisms cannot be completely dismissed, though. One possibility is that the barbiturates bind and dissociate on the microsecond time scale and induce a conformational change to a new closed state of the channel. In this scenario, the transition rates between the open and new closed states determine bursting and relaxation kinetics. These rates would have to be exquisitely sensitive to the difference in chemical structure between PB and Barb to account for the 100-fold difference in the kinetic actions of these drugs.

The question of the location of the barbiturate binding site(s) might be answered more convincingly by site-directed mutagenesis experiments as has been done for open channel blockers such as QX-222 (Charnet et al., 1990). Yost and Dodson (1993) have argued that the site of action for amobarbital does not involve amino acids at the 10′ level (near the center of the membrane) of the M2 transmembrane region of the channel. This does not, however, rule out other sites within the pore of the channel, nor does it rule out the 10′ level as being part of the binding site for other barbiturates. Inhibition of AMPA-selective glutamate receptor channels by PB is influenced by amino acids within the M2 region of the channel pore (Yamakura et al., 1995). However, so far there is no kinetic evidence that PB acts as blocker of this channel.

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REFERENCES

Auerbach, A. 1993. A statistical analysis of acetylcholine receptor activation in *Xenopus* myocytes: stepwise versus concerted models of gating. *J. Physiol. (Lond.).* 461:339–378.

Barann, M., M. Gothert, K. Fink, and H. Bonisch. 1993. Inhibition by anaesthetics of [3H]-guanidinium flux through the voltage-gated sodium channel and the cation channel of the 5-HT3 receptor of NIE-115 neuroblastoma cells. *Naunyn-Schmiedebergs. Arch. Pharmacol.* 347:125–132.

Charnet, P., C. Labarca, R.J. Leonard, N.J. Vogelaar, L. Czyzyk, A. Gouin, N. Davidson, and H.A. Lester. 1990. An open-channel blocker interacts with adjacent turns of α-helices in the nicotinic acetylcholine receptor. *Nature* 347:87–95.

Colquhoun, D., and A.G. Hawkes. 1995a. The principles of the stochastic interpretation of ion-channel mechanisms. In *Single Channel Recording.* B. Sakmann and E. Neher, editors. Plenum Press, New York. 397–482.

Colquhoun, D., and A.G. Hawkes. 1995b. A Q-matrix cookbook. In *Single Channel Recording.* B. Sakmann and E. Neher, editors. Plenum Press, New York. 589–633.

deArmendi, A.J., P.H. Tonner, B. Bugge, and K.W. Miller. 1993. Barbital uration is dependent on the conformational state of the acetylcholine receptor. *Anesthesiology. 79*:1035–1041.

Dilger, J.P., and R.S. Brett. 1990. Direct measurement of the concentration- and time-dependent open probability of the nicotinic acetylcholine receptor channel. *Biophys. J.* 57:729–731.

Dilger, J.P., and Y. Liu. 1992. Desensitization of acetylcholine receptors in BC3H-1 cells. *Pflüg. Arch.* 420:479–485.

Dilger, J.P., R.S. Brett, and L. Lesko. 1992. Effects of iso-1urane on acetylcholine receptor channels. 1. Single-channel currents. *Mol. Pharmacol.* 41:127–133.

Dilger, J.P., A.M. Vidal, H.I. Mody, and Y. Liu. 1994. Evidence for direct actions of general anaesthetics on an ion channel protein: a new look at a unified mechanism of action. *Anesthesiology. 81*:431–442.

Dodson, B.A., R.R. Urh, and K.W. Miller. 1990. Relative potencies for barbiturate binding to the *Torpedo* acetylcholine receptor. *Br. J. Pharmacol.* 101:710–714.

French-Mullen, J.M.H., J.L. Barker, and M.A. Rogawski. 1993. Calcium current block by (−)-pentobarbital, phenobarbital, and CHEB but not (+)-pentobarbital in acutely isolated hippocampal CA1 neurons: comparison with effects on GABA-activated Cl current. *J. Neurosci.* 13:3211–3221.

Firestone, L.L., J.C. Miller, and K.W. Miller. 1986a. Tables of physical and pharmacological properties of anaesthetics. In *Molecular and Cellular Mechanisms of Anaesthetics.* S.H. Roth and K.W. Miller, editors. Plenum Publishing Corp., New York. 267–277.

Firestone, L.L., J.F. Saucier, L.M. Braswell, and K.W. Miller. 1986b. Actions of general anaesthetics on acetylcholine receptor-rich membranes from *Torpedo californica.* *Anesthesiology.* 64:694–702.

Firestone, L.L., J.K. Allinmof, and K.W. Miller. 1994. Does general anesthetic-induced desensitization of the Torpedo acetylcholine receptor correlate with lipid disordering? *Mol. Pharmacol.* 46:508–515.

Fragen, R.J. 1994. Clinical pharmacology and applications of intravenous anesthetic induction agents. In *The Pharmacologic Basis of Anesthesiology.* T.A. Bowdle, A. Horita, and E.D. Kharasch, editors. Churchill Livingstone, New York. 319–336.

Frenkel, C., D.S. Duch, and B.W. Urban. 1990. Molecular actions of pentobarbital isomers on sodium channels from human brain cortex. *Anesthesiology.* 72:640–649.

Gage, P.W., and D. McKinnon. 1985. Effects of pentobarbitone on acetylcholine-activated channels in mammalian muscle. *Br. J. Pharmacol.* 85:229–235.

Hall, A.C., W.R. Lieb, and N.P. Franks. 1994. Insensitivity of P-type calcium channels to inhalational and intravenous general anaesthetics. *Anesthesiology. 81*:117–123.

Hamill, O.P., A. Marty, E. Neher, B. Sakmann, and F.J. Sigworth. 1981. Improved patch-clamp techniques for high-resolution current recording from cells and cell free membrane patches. *Pflüg. Arch.* 391:85–100.

Jacobson, L., G. Pocock, and C.D. Richards. 1991. Effects of pentobarbital on the properties of nicotinic channels of chromaffin cells. *Eur. J. Pharmacol.* 202:331–339.

Lee-Son, S., B.E. Waud, and D.R. Waud. 1975. A comparison of the potencies of a series of barbiturates at the neuromuscular junction and on the central nervous system. *J. Pharmacol. Exp. Therap.* 195:251–256.

Liu, Y., and J.P. Dilger. 1991. Opening rate of acetylcholine receptor channels. *Biophys. J.* 60:424–432.

Marszałek, W., and T. Narahashi. 1993. Use-dependent pentobarbital block of kainate and quisqualate currents. *Brain Res.* 608:7–15.

Murrell, R.D., M.S. Braun, and D.A. Haydon. 1991. Actions of n-alcohols on nicotinic acetylcholine receptor channels in cultured rat myotubes. *J. Physiol. (Lond.).* 437:431–448.

Neher, E. 1983. The charge carried by single-channel currents of rat cultured muscle cells in the presence of local anaesthetics. *J. Physiol. (Lond.).* 339:663–678.

Roth, S., S.A. Forman, L.M. Braswell, and K.W. Miller. 1989. Actions of pentobarbital enantiomers on nicotinic cholinergic receptors. *Mol. Pharmacol.* 36:874–880.

Sine, S.M., and J.H. Steinbach. 1984. Activation of a nicotinic acetylcholine receptor. *Biophys. J.* 45:175–185.

Tanelian, D.L., P. Kosek, I. Mody, and M.B. MacIver. 1993. The role of the GABA A receptor/chloride channel complex in anesthesia. *Anesthesiology.* 78:757–776.

Yamakura, T., K. Sakimura, M. Mishina, and K. Shimoji. 1995. The sensitivity of AMPA-selective glutamate receptor channels to pentobarbital is determined by a single amino acid residue of the α2 subunit. *FEBS Lett.* 374:412–414.

Yellen, G. 1984. Ionic permeation and blockade in Ca2+-activated K+ channels of bovine chromaffin cells. *J. Gen. Physiol.* 84:157–186.

Yost, C.S., and B.A. Dodson. 1993. Inhibition of the nicotinic acetylcholine receptor by barbiturates and by procaine: do they act at different sites? *Cell. Mol. Neurobiol.* 13:159–172.