# Spike-and-Wave Oscillations Based on the Properties of GABA<sub>B</sub> Receptors

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Neocortical and thalamic neurons are involved in the genesis of generalized spike-and-wave (SW) epileptic seizures. The cellular mechanism of SW involves complex interactions between intrinsic neuronal firing properties and multiple types of synaptic receptors, but because of the complexity of these interactions the exact details of this mechanism are unclear. In this paper these types of interactions were investigated by using biophysical models of thalamic and cortical neurons. It is shown first that, because of the particular activation properties of GABA<sub>B</sub> receptor-mediated responses, simulated field potentials can display SW waveforms if cortical pyramidal cells and interneurons generate prolonged discharges in synchrony, without any other assumptions. Here the "spike" component coincided with the synchronous firing, whereas the "wave" component was generated mostly by slow GABA<sub>B</sub>-mediated K<sup>+</sup> currents. Second, the model suggests that intact thalamic circuits can be forced into a  $\sim$ 3 Hz oscillatory mode by corti-

Generalized spike-and-wave (SW) patterns characterize the human electroencephalogram during several types of epilepsy as well as in animal models of absence seizures. Initially suggested by Jasper and Kershman (1941), the possible involvement of the thalamus in SW seizures was shown by recordings of thalamic nuclei in humans during absence attacks (Williams, 1953; Prevett et al., 1995). An important role for the thalamus also is supported by electrophysiological recordings in experimental models of SW seizures, which show that cortical and thalamic cells fire prolonged discharges in phase with the "spike" component, whereas the "wave" is characterized by a silence in all cell types (Pollen, 1964; Steriade, 1974; Avoli et al., 1983; McLachlan et al., 1984; Buzsáki et al., 1990; Inoue et al., 1993). Electrophysiological recordings also indicate that spindle oscillations, which are generated by thalamic circuits (Steriade et al., 1990, 1993), can be transformed gradually into SW discharges, and all manipulations that promote or antagonize spindles have the same effect on SW (Kostopoulos et al., 1981a,b; McLachlan et al., 1984). Finally, it has been demonstrated that SW patterns disappear after thalamic lesions or after inactivation of the thalamus (Pellegrini et al., 1979; Avoli and Gloor, 1981; Vergnes and Marescaux, 1992).

A series of pharmacological results suggests that  $GABA_B$  receptors play a critical role in the genesis of SW discharges in rats, because  $GABA_B$  agonists exacerbate seizures whereas  $GABA_B$  antagonists suppress them (Hosford et al., 1992; Snead, 1992;

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This research was supported by the Medical Research Council of Canada (MT-13724). We thank D. Contreras and N. Kopell for comments on this manuscript. Correspondence should be addressed to Dr. Alain Destexhe at the above address. Copyright © 1998 Society for Neuroscience 0270-6474/98/189099-13\$05.00/0 cothalamic feedback. Here again, this property was attributable to the characteristics of  $\mathsf{GABA}_{\mathsf{B}}\text{-mediated}$  inhibition. Third, in the thalamocortical system this property can lead to generalized  $\sim 3$  Hz oscillations with SW field potentials. The oscillation consisted of a synchronous prolonged firing in all cell types, interleaved with a  $\sim 300$  msec period of neuronal silence, similar to experimental observations during SW seizures. This model suggests that SW oscillations can arise from thalamocortical loops in which the corticothalamic feedback indirectly evokes  $\mathsf{GABA}_{\mathsf{B}}\text{-mediated}$  inhibition in the thalamus. This mechanism is shown to be consistent with a number of different experimental models, and experiments are suggested to test its consistency.

Key words: computational models; thalamus; cerebral cortex; epilepsy; absence; intrinsic properties; low-threshold spikes; spindle oscillations; thalamocortical

Puigcerver et al., 1996; Smith and Fisher, 1996). The anti-absence drug clonazepam seems to act by diminishing  $GABA_B$ -mediated IPSPs in thalamocortical (TC) cells, reducing their tendency to burst in synchrony (Huguenard and Prince, 1994a; Gibbs et al., 1996). In ferret thalamic slices, spindle oscillations can be transformed into slower ~3 Hz oscillations after blocking  $GABA_A$  receptors, and, like SW, these oscillations are suppressed by  $GABA_B$  receptor antagonists (von Krosigk et al., 1993). These experiments could be replicated by computational models of thalamic circuits (Destexhe and Sejnowski, 1995; Destexhe et al., 1996a; Golomb et al., 1996).

Although these results could suggest a thalamic origin of SW seizures involving  $GABA_B$ -mediated mechanisms, clear evidence suggests a determinant role for the cortex: thalamic injections of high doses of  $GABA_A$  antagonists such as penicillin (Ralston and Ajmone-Marsan, 1956; Gloor et al., 1977) or bicuculline (Steriade and Contreras, 1998) led to 3–4 Hz oscillations, with no sign of SW discharge. On the other hand, injection of the same drugs to the cortex, with no change in the thalamus, resulted in seizure activity with SW patterns (Gloor et al., 1977; Steriade and Contreras, 1998).

These experiments show that both cortical and thalamic neurons are necessary to generate SW rhythms and that both  $GABA_A$  and  $GABA_B$  receptors are actively involved, but the exact mechanisms are unclear. In this paper a thalamocortical loop mechanism for the genesis of SW oscillatory patterns was investigated by the use of computational models that were based on the complex intrinsic firing properties of thalamic and cortical neurons (see Llinás, 1988) and the properties particular to each receptor type (Destexhe et al., 1998b).

#### MATERIALS AND METHODS

All models that are shown here were based on biophysical representations of the ionic mechanisms underlying synaptic currents, field potential generation, intrinsic firing properties, and network behavior. The modeling methods that were used to simulate these various aspects are described successively.

Synaptic currents. Postsynaptic currents mediated by glutamate AMPA and NMDA receptors as well as by GABAergic GABA<sub>A</sub> and GABA<sub>B</sub> receptors were simulated by kinetic models of postsynaptic receptors (Destexhe et al., 1994, 1998b). When a spike occurred in the presynaptic cell, a brief pulse of transmitter concentration (0.5 mM during 0.3 msec) was simulated in the synaptic cleft, and binding of the transmitter to postsynaptic receptors occurred according to simple open/closed kinetics, leading to a transient increase of the postsynaptic current described by the following equation (Destexhe et al., 1994):

$$I_{\rm syn} = \bar{g}_{\rm syn} \, m \, \left( V - E_{\rm syn} \right) \, \text{and} \tag{1}$$

$$\frac{dm}{dt} = \alpha \left[T\right] (1-m) - \beta m, \qquad (2)$$

where  $I_{\rm syn}$  is the postsynaptic current,  $\neg g_{\rm syn}$  is the maximal conductance, *m* is the fraction of open receptors,  $E_{\rm syn}$  is the reversal potential, [*T*] is the transmitter concentration in the cleft, and  $\alpha$  and  $\beta$  are forward and backward binding rate constants of *T* to open the receptors. This scheme was used to simulate AMPA, NMDA, and GABA<sub>A</sub> types of receptors, with the following parameters:  $E_{\rm syn} = 0$  mV,  $\alpha = 0.94 \times 10^6 M^{-1} s^{-1}$ ,  $\beta = 180 s^{-1}$  for AMPA receptors;  $E_{\rm syn} = 0$  mV,  $\alpha = 11 \times 10^4 M^{-1} s^{-1}$ ,  $\beta = 6.6 s^{-1}$  for NMDA receptors; and  $E_{\rm syn} = -80$  mV,  $\alpha = 20 \times 10^6$  $M^{-1} s^{-1}$ ,  $\beta = 160 s^{-1}$  for GABA<sub>A</sub> receptors. These parameters were obtained by fitting the model to postsynaptic currents recorded experimentally (see Destexhe et al., 1998b). In addition, NMDA receptors had a voltage-dependent term corresponding to an extracellular Mg<sup>2+</sup> concentration of 2 mm [Jahr and Stevens (1990); see Destexhe et al. (1998b) for the details of implementation].

The modeling of slow GABA<sub>B</sub> receptor-mediated inhibition required a more complex scheme to capture the nonlinear properties of this type of interaction (Destexhe and Sejnowski, 1995). The activation properties of GABA<sub>B</sub> receptors were based on the following steps: (1) the binding of GABA on the GABA<sub>B</sub> receptor, leading to the activated receptor; (2) the activated GABA<sub>B</sub> receptor catalyzes the activation of G-proteins in the intracellular side; (3) the binding of activated G-proteins to open K<sup>+</sup> channels. These steps are described by the following equations:

$$I_{\text{GABAB}} = \bar{g}_{\text{GABAB}} \frac{s^n}{s^n + K_{\text{D}}} (V - E_{\text{K}}), \qquad (3)$$

$$\frac{dr}{dt} = K_1 [T] (1 - r) - K_2 r$$
, and (4)

$$\frac{ds}{dt} = K_3 r - K_4 s, \tag{5}$$

where [T] is the GABA concentration in the synaptic cleft, r is the fraction of GABA<sub>B</sub> receptors in the activated form, s is the normalized G-protein concentration in activated form,  $\bar{g}_{\text{GABA}_{\text{B}}}$  is the maximal postsynaptic conductance of K<sup>+</sup> channels,  $K_{\text{D}}$  is the dissociation constant of G-protein binding on K<sup>+</sup> channels, V is the postsynaptic membrane potential, and  $E_{\text{K}}$  is the equilibrium potential for K<sup>+</sup>. The fitting of this model to experimental GABA<sub>B</sub> responses led to the following values of parameters (Destexhe et al., 1998b):  $K_{\text{D}} = 100$ ,  $K_1 = 9 \times 10^4 M^{-1} s^{-1}$ ,  $K_2 = 1.2 s^{-1}$ ,  $K_3 = 180 s^{-1}$ , and  $K_4 = 34 s^{-1}$ , with n = 4 binding sites.

*Field potentials.* Extracellular field potentials were calculated from postsynaptic currents in single-compartment models according to the model of Nunez (1981):

$$V_{\rm ext} = \frac{R_{\rm e}}{4\pi} \sum_{\rm j} \frac{I_{\rm j}}{r_{\rm j}},\tag{6}$$

where  $V_{\text{ext}}$  is the electrical potential at a given extracellular site,  $R_{\text{e}} = 230$  $\Omega$ cm is the extracellular resistivity,  $I_{\text{j}}$  is the postsynaptic current, and  $r_{\text{j}}$  is the distance between the site of generation of  $I_{\text{i}}$  and the extracellular site.

Field potentials were calculated from a single cell receiving 200 simulated synapses (100 excitatory synapses had AMPA and NMDA receptor types, and 100 inhibitory synapses had  $GABA_A$  and  $GABA_B$  receptor types.

tors; see the scheme in Fig. 1*B*). In this case, trains of presynaptic action potentials were generated individually for each synapse. To avoid possible artifactual effects because of the coincident timing of action potentials at different synapses, a random time jitter of  $\pm 1$  msec was included in the timing of each presynaptic action potential.

*Intrinsic currents.* Intrinsic voltage-dependent or calcium-dependent currents were modeled by kinetic models of the Hodgkin and Huxley (1952) type. These intrinsic membrane currents were described by the following generic equation:

$$I_{\rm int} = \bar{g}_{\rm int} \, m^{\rm N} h^{\rm M} \, (V - E_{\rm int}), \tag{7}$$

$$\frac{dm}{dt} = \alpha_{\rm m} \left( 1 - m \right) - \beta_{\rm m} \, m, \, \text{and} \tag{8}$$

$$\frac{dh}{dt} = \alpha_{\rm h} \left( 1 - h \right) - \beta_{\rm h} h, \tag{9}$$

where  $I_{int}$  is the intrinsic membrane current,  $\neg g_{int}$  is the maximal conductance, and  $E_{int}$  is the reversal potential. The gating properties of the current were dependent on N activation gates and M inactivation gates, with m and h representing the fraction of gates in open form, and with respective rate constants  $\alpha_m$ ,  $\beta_m$ ,  $\alpha_h$ , and  $\beta_h$ . Rate constants were dependent on either membrane voltage (V) or intracellular calcium concentration.

Thalamocortical networks. Network models were based on singlecompartment representations of thalamic and cortical neurons. The thalamocortical network was simulated with four cell types: cortical pyramidal cells (PY), cortical interneurons (IN), thalamic reticular cells (RE), and thalamocortical (TC) cells. Cortical cells represent layer VI of the cerebral cortex, in which PY cells constitute the major source of corticothalamic fibers. Because corticothalamic PY cells receive a significant proportion of their excitatory synapses from ascending thalamic axons (Hersch and White, 1981; White and Hersch, 1982), these cells mediate a monosynaptic excitatory feedback loop (thalamus–cortex– thalamus) that has been modeled here. Each layer of cells has been arranged in one dimension (connectivity is schematized in Fig. 4A). This one-dimensional network model with four cell types is a greatly simplified representation of the multilayered structure of the thalamocortical system, but no additional complexity was required.

The cellular models had intrinsic and synaptic currents described by the membrane equation:

$$C_{\rm m} \dot{V}_{\rm i} = -g_{\rm L} \left( V_{\rm i} - E_{\rm L} \right) - \sum_{j} I_{\rm int}^{\rm ji} - \sum_{k} I_{\rm syn}^{\rm ki},$$
 (10)

where  $V_i$  is the membrane potential,  $C_m = 1 \ \mu F/cm^2$  is the specific capacity of the membrane,  $g_L$  is the leakage conductance, and  $E_L$  is the leakage reversal potential. Intrinsic and synaptic currents are represented by  $I_{int}^{li}$  and  $I_{syn}^{ki}$ , respectively.

by  $I_{int}^{ii}$  and  $I_{syn}^{ki}$ , respectively. The synaptic currents  $I_{syn}^{ki}$  from presynaptic cell k to postsynaptic cell i, were simulated by activating a short pulse of transmitter when cell k fired an action potential (see above). The receptor types present in synaptic connections between cells depended on the cell type. All excitatory connections (TC $\rightarrow$ RE, TC $\rightarrow$ IN, TC $\rightarrow$ PY, PY $\rightarrow$ PY, PY $\rightarrow$ IN,  $PY \rightarrow RE, PY \rightarrow TC$ ) were mediated by AMPA receptors; some inhibitory connections (RE $\rightarrow$ TC, IN $\rightarrow$ PY) were mediated by a mixture of GABA<sub>A</sub> and GABA<sub>B</sub> receptors, whereas intra-RE connections were mediated by GABA<sub>A</sub> receptors. Simulations also were performed with NMDA receptors added to all excitatory connections (with maximal conductance set to 25% of that of AMPA), and no appreciable difference was observed. They therefore were not included in the present figures. The total synaptic conductance on each neuron was the same for cells of the same type and was expressed as the sum over all individual synaptic conductances of the same connection type. The total conductances corresponding to the reference state, displaying spindle oscillations, were 0.2  $\mu$ S (ÅMPA, TC $\rightarrow$ RE), 0.2  $\mu$ S (GABA<sub>A</sub>, RE $\rightarrow$ RE), 0.02  $\mu$ S (GABA<sub>A</sub>, RE $\rightarrow$ TC), 0.04  $\mu$ S (GABA<sub>B</sub>, RE $\rightarrow$ TC), 0.6  $\mu$ S (AMPA, PY $\rightarrow$ PY), 0.2  $\mu$ S (AMPA, PY $\rightarrow$ IN), 0.15  $\mu$ S (GABA<sub>A</sub>, IN $\rightarrow$ PY), 0.03  $\mu$ S (GABA<sub>B</sub>, IN $\rightarrow$ PY), 1.2  $\mu$ S (AMPA, PY $\rightarrow$ RE), 0.01  $\mu$ S (AMPA, PY $\rightarrow$ TC), 1.2  $\mu$ S (AMPA, TC $\rightarrow$ PY), and 0.4  $\mu$ S (AMPA, TC $\rightarrow$ IN).

The connectivity between thalamic and cortical layers was topographic: within the thalamus and within cortex, each axon contacted the 11 nearest neighbors to the presynaptic cell. The axonal divergence was of 21 cells for projections between thalamus and cortex. The connection topology, values of synaptic conductances, and robustness of the network were described in detail in a previous study (Destexhe et al., 1998a).

All intrinsic membrane currents  $I_{int}^{ii}$  were described by a variant of the Hodgkin and Huxley (1952) model (Eqs. 7-9). All cell types had Na and K<sup>+</sup> currents for generating action potentials, for which the kinetics was taken from Traub and Miles (1991). Additional currents conferred to each cell type the most salient features of its intrinsic firing patterns. Thalamic cells produced bursts of action potentials because of the presence of a T-current (see *inset* in Fig. 3A). In TC cells, in addition to  $I_{\rm T}$ , the presence of  $I_{\rm h}$  conferred oscillatory properties. The upregulation of  $I_{\rm h}$  by intracellular Ca<sup>2+</sup> led to waxing and waning properties of these oscillations, as detailed in previous models (Destexhe et al., 1993, 1996a, 1998a). In RE cells the T-current was of slower kinetics, as modeled previously (Destexhe et al., 1996b). Models for cortical cells were kept as simple as possible to reproduce their repetitive firing properties (see inset in Fig. 4A). IN cells contained no other current than was necessary for action potentials, producing similar firing patterns to "fast-spiking" cells (Connors and Gutnick, 1990). PY cells had one additional slow voltagedependent  $K^+$  current  $(I_M)$  generating adapting trains of action potentials, similar to "regular-spiking" pyramidal cells (Connors and Gutnick, 1990). The conductance values and the activation properties of all intrinsic membrane currents were identical to a previous study (Destexhe et al., 1998a).

Field potentials were calculated from network simulations. In this case only cortical pyramidal cells were considered and were arranged equidistantly in one dimension (intercellular distance of 20  $\mu$ m). Then field potentials at a given extracellular site were calculated from postsynaptic currents:

$$V_{\rm ext} = \frac{R_{\rm e}}{4\pi} \sum_{\rm i,k} \frac{I_{\rm syn}^{\rm ki}}{r_{\rm i}},\tag{11}$$

where  $r_i$  is the distance between each PY cell and the extracellular site. In some cases the contribution of the voltage-dependent current  $I_M$  in

field potentials was evaluated according to the relation:

$$V_{\text{ext}} = \frac{R_{\text{e}}}{4\pi} \sum_{i} \frac{(I_{\text{M}}^{\text{i}} + \Sigma_{k} I_{\text{syn}}^{\text{ki}})}{r_{\text{i}}}, \qquad (12)$$

where  $I_{\rm M}^i$  is the voltage-dependent K<sup>+</sup> current responsible for adaptation of repetitive firing in the *i*th PY cells.

All models were simulated by using NEURON (Hines and Carnevale, 1997) and were run on a Sparc-20 workstation (Sun Microsystems, Mountain View, CA).

### RESULTS

The mechanism underlying a slow oscillation similar to SW is explained in three steps: (1) the nonlinear activation properties of GABA<sub>B</sub> responses can lead to the generation of SW waveforms in field potentials; (2) intact thalamic circuits can be forced into a  $\sim$ 3 Hz oscillation by corticothalamic feedback; (3) the combination of these two factors can generate  $\sim$ 3 Hz oscillations with SW field potentials in thalamocortical networks. These points are considered successively.

## The nonlinear activation properties of GABA<sub>B</sub> responses

A property consistently observed for  $GABA_B$  responses is that they require high stimulus intensities to be evoked, as shown in hippocampal (Dutar and Nicoll, 1988; Davies et al., 1990) and thalamic slices (Kim et al., 1997). This property can be reproduced under certain nonlinearity assumptions in the G-protein transduction mechanisms evoked by  $GABA_B$  receptors; assuming that the binding of four G-proteins is required to activate K<sup>+</sup> channels is enough to provide a nonlinear stimulus dependence similar to  $GABA_B$  responses (Destexhe and Sejnowski, 1995). The multiplicity of binding sites of G-proteins is indeed in agreement with the tetrameric structure of K<sup>+</sup> channels (Hille, 1992) and the cooperativity evidenced in the activation of  $GABA_B$  responses (Sodickson and Bean, 1996).

The nonlinear stimulus dependence in the model of  $GABA_B$  currents is illustrated in Figure 1*A*. An isolated presynaptic spike could not evoke detectable  $GABA_B$  current (Fig. 1*A1*), in agreement with the absence of  $GABA_B$ -mediated miniature events (Otis and Mody, 1992; Thompson and Gahwiler, 1992; Thompson, 1994). However, a burst of 5–10 high-frequency spikes is a very powerful means of evoking  $GABA_B$  responses (Fig. 1*A2*). The latter feature is consistent with the observation that  $GABA_B$  responses appear only under high-intensity stimulus conditions (Dutar and Nicoll, 1988; Davies et al., 1990) and the evidence that bursts of high-frequency action potentials are an ideal presynaptic signal to evoke  $GABA_B$  currents (Huguenard and Prince, 1994b; Kim et al., 1997). In the model this property is obtained from the fact that a sufficient level of G-proteins must be accumulated to evoke significant K<sup>+</sup> current.

### Possible role of GABA<sub>B</sub>-mediated currents in generating spike-and-wave field potentials

The possible role of the particular activation properties of GABA<sub>B</sub> currents in generating SW patterns was investigated by simulating field potentials from the postsynaptic currents generated by 100 excitatory synapses (AMPA and NMDA receptors) and 100 inhibitory synapses (GABA<sub>A</sub> and GABA<sub>B</sub> receptors; see scheme in Fig. 1B and Materials and Methods). With presynaptic trains consisting of single spikes, the voltage showed mixed EPSP/ IPSP sequences, and the field potential was dominated by negative deflections (Fig. 1C1). By contrast, bursts of high-frequency presynaptic spikes produced mixed EPSP/IPSPs, followed by large GABA<sub>B</sub>-mediated IPSPs in the cell (Fig. 1C2). In this case the fast EPSP/IPSPs generated spiky field potentials, followed by a slow positive wave caused by GABA<sub>B</sub> currents. This simple simulation therefore shows that, if excitatory and inhibitory cells generate high-frequency discharges in synchrony and if GABA<sub>B</sub> receptors are present, sufficient conditions are brought together to generate field potential waveforms consisting of interleaved spikes and waves.

The effect of various parameters on the morphology of simulated SW complexes was investigated in Figure 2*A*. When excitatory synapses discharged earlier than inhibitory synapses (2 and 5 msec latency), the spike component was enhanced. Spike and wave components also were influenced by synaptic conductances. AMPA and NMDA conductances affected primarily the negative peak of the spike component (Fig. 2*B*, top trace), whereas the positive peak was influenced mostly by GABA<sub>A</sub> conductances (Fig. 2*B*, middle trace). GABA<sub>B</sub> conductances had few effects on the spike component but mostly affected the wave (Fig. 2*B*, bottom trace).

### Intact thalamic circuits can be forced into $\sim$ 3 Hz oscillations because of GABA<sub>B</sub>-mediated currents

To investigate how this type of field potentials can be generated by the thalamocortical system, we first turn to the behavior of thalamic circuits, and, more particularly, we turn to how they are controlled by the cortex. An important behavior of thalamic networks is their propensity to generate oscillations such as the 7–14 Hz spindle oscillations (Steriade et al., 1993; von Krosigk et al., 1993). Although these oscillations are generated in the thalamus, the neocortex has been shown to trigger them powerfully (Steriade et al., 1972; Roy et al., 1984; Contreras and Steriade, 1996), and the corticothalamic feedback has been shown to exert a decisive control over thalamic oscillations (Contreras et al., 1996). Figure 1. Simulation of spike and wave in field potentials, based on the properties of  $GABA_B$  receptors. A, Nonlinear activation properties of GABA<sub>B</sub> receptors. If the binding of four G-proteins is needed to activate the K<sup>+</sup> channels associated with GABA<sub>B</sub> receptors, then GABA<sub>B</sub>-mediated inhibitory responses are dependent on the number of presynaptic action potentials. A1, With a single presynaptic spike no GABA<sub>B</sub> response was detectable. A2, A burst of presynaptic spikes led to sufficient accumulation of G-proteins to generate the slow IPSP. B, Scheme for the model of local field potentials. Excitatory and inhibitory presynaptic trains of action potentials were generated to stimulate various postsynaptic receptor types (AMPA, NMDA, GABAA, and GABA<sub>B</sub>). One hundred synapses of each type were simulated, and the synaptic currents were integrated into a single compartment model and used to calculate the extracellular field potential at a distance of 5  $\mu$ m from the simulated neuron. C, Field potentials generated by single spikes and bursts of spikes. With single spikes (C1) the mixed EPSP/IPSP sequence led to negative deflections in the field potentials. With bursts of spikes (C2) the fast spiky components alternate with slow positive deflections, similar to spike-and-wave patterns. These slow positive waves are attributable to the activation of GABA<sub>B</sub>-mediated currents (arrows). Conductance values are 4, 1, 1.5, and 4 nS for individual AMPA, NMDA, GABA<sub>A</sub>, and GABA<sub>B</sub> synapses, respectively.

In computational models, reproducing this cortical control required more powerful corticothalamic EPSPs on RE cells as compared with TC cells (Destexhe et al., 1998a). In these conditions the excitation of corticothalamic cells led to mixed EPSPs and IPSPs in TC cells in which the IPSP was dominant, consistent with experimental observations (Burke and Sefton, 1966; Deschênes and Hu, 1990). If cortical EPSPs and IPSPs from RE cells were of comparable conductance, cortical feedback could not evoke oscillations in the thalamic circuit because of shunting effects between EPSPs and IPSPs (Destexhe et al., 1998a). The most likely reason for these experimental and modeling evidences for "IPSP dominance" in TC cells is that RE cells are extremely sensitive to cortical EPSPs (Contreras et al., 1993), probably because of a powerful T-current in dendrites (Destexhe et al., 1996b). In addition, cortical synapses contact only the distal dendrites of TC cells (Liu et al., 1995) and probably are attenuated for this reason. Taken together, these data suggest that corticothalamic feedback operates mainly by eliciting bursts in RE cells, which in turn evoke powerful IPSPs on TC cells that in large part overwhelm the direct cortical EPSPs.

The effect of corticothalamic feedback on the thalamic circuit is depicted in Figure 3*A*: simulated cortical EPSPs evoked bursts in RE cells (Fig. 3*B, arrow*), which recruited TC cells via IPSPs, and triggered a  $\sim 10$  Hz oscillation in the circuit. During the oscillation TC cells rebounded after GABA<sub>A</sub>-mediated IPSPs once every two cycles, and RE cells discharged only a few spikes,



evoking GABA<sub>A</sub>-mediated IPSPs in TC cells with no significant GABA<sub>B</sub> currents (Fig. 3*B*). These features are typical of spindle oscillations (Steriade et al., 1993; von Krosigk et al., 1993).

Repetitive stimulation of the same thalamic circuit at 3 Hz with larger intensity (14 spikes every 333 msec) entrained the system into a different type of oscillatory behavior (Fig. 3*C*). All cell types were entrained to discharge in synchrony at  $\sim$ 3 Hz. On the other hand, repetitive stimulation at 3 Hz with low intensity produced spindle oscillations (Fig. 3*D*) similar to those in Figure 3*A*. Strong-intensity stimulation at 10 Hz led to quiescence in TC cells (Fig. 3*E*) because of sustained GABA<sub>B</sub> currents, similar to a previous analysis [Lytton et al. (1997), their Fig. 12].

These simulations indicate that strong corticothalamic feedback at 3 Hz can force thalamic circuits in a different type of oscillation. Cortical EPSPs force RE cells to fire large bursts (Fig. 3*C*, *arrows*), fulfilling the conditions needed to activate GABA<sub>B</sub> responses (see Fig. 1*A*). The consequence is that TC cells were "clamped" at hyperpolarized levels by GABA<sub>B</sub> IPSPs during ~300 msec before they could rebound. The nonlinear properties of GABA<sub>B</sub> responses are therefore responsible for the coexistence between two types of oscillations in the same circuit: mild corticothalamic feedback recruits the circuit in ~10 Hz spindle oscillations, whereas strong feedback at 3 Hz could force the intact circuit at the same frequency because of the nonlinear activation properties of intrathalamic GABA<sub>B</sub> responses.



*Figure 2.* Factors determining the morphology of simulated spike-andwave complexes. *A*, Effect of a latency between the firing of excitatory and inhibitory synapses. *Control*, Simulation is identical to Figure 1C2. If excitatory synapses discharged earlier than inhibitory synapses (2 and 5 msec latency), the spike component was enhanced. *B*, Effect of synaptic receptor types. The 5 *msec latency* simulation from *A* was repeated here with different values for synaptic conductances. A 90% reduction of AMPA/NMDA conductances (*top trace*), GABA<sub>A</sub> conductances (*middle trace*), or GABA<sub>B</sub> conductances (*bottom trace*) affected the morphology of the SW patterns. The *right column* shows the last complexes at higher resolution.

# Suppression of intrathalamic $GABA_A$ -mediated inhibition does not generate spike and wave

The impact of this mechanism at the network level was explored using a thalamocortical network consisting in different layers of cortical and thalamic cells (see details in Materials and Methods). The network included thalamic TC and RE cells and a simplified representation of the deep layers of the cortex with pyramidal cells and interneurons (Fig. 4*A*). In control conditions (Fig. 4*B*) the network generated synchronized spindle oscillations with cellular discharges in phase between in all cell types, as observed experimentally (Contreras and Steriade, 1996). TC cells discharged on average once every two cycles after GABA<sub>A</sub>-mediated IPSPs, whereas all other cell types discharged approximately at every cycle at  $\sim 10$  Hz, consistent with the typical features of spindle oscillations observed intracellularly (Steriade et al., 1990; von Krosigk et al., 1993). The simulated field potentials displayed successive negative deflections at  $\sim 10$  Hz (Fig. 4*B*;

in agreement with the pattern of field potentials during spindle oscillations) (Steriade et al., 1990). Consistent with the analysis of Figure 1*C1*, this pattern of field potentials was generated by the limited discharge in PY cells, which fired approximately one spike per oscillation cycle.

When GABA<sub>A</sub> receptors were suppressed in thalamic cells in this model, with cortical inhibition intact, spindle oscillations were transformed into slower oscillation patterns at 3-5 Hz (Fig. 4C). In this case there was an increase in synchrony, as indicated by the TC cells that fired at every cycle of the oscillation. RE cells generated prolonged burst discharges, leading to GABA<sub>B</sub>mediated IPSPs in TC cells and, consequently, to a slow oscillation frequency. The field potentials consisted of successive negative deflections (Fig. 4C, bottom) similar to that of spindles. This pattern of field potentials was generated by PY cells that discharged approximately single spikes at each cycle of the oscillation (similar to Fig. 1C1). This simulation therefore suggests that removing intrathalamic GABA<sub>A</sub>-mediated inhibition affects the oscillation frequency but does not generate SW, because pyramidal cells are still under the strict control of cortical fast inhibition. This is in agreement with in vivo injections of bicuculline into the thalamus, which reported slow oscillations with increased thalamic synchrony, but no SW patterns in the field potentials (Ralston and Ajmone-Marsan, 1956; Steriade and Contreras, 1998).

# Suppression of intracortical GABA<sub>A</sub>-mediated inhibition leads to spike and wave

On the other hand, the alteration of GABA<sub>A</sub> receptors in the cortex had a considerable impact in generating SW. When GABA<sub>A</sub>-mediated inhibition was reduced in the cortex, with no change in thalamic inhibitory mechanisms, then spindle oscillations transformed into 2–3 Hz SW-like discharges (Fig. 5). With intracortical fast inhibition decreased by 50%, increased occurrences of prolonged high-frequency discharges were seen during spindle oscillations (Fig. 5*A*). In field potentials these events tended to generate large-amplitude negative deflections, followed by small-amplitude positive waves (Fig. 5*A*, *bottom*).

With totally suppressed GABA<sub>A</sub>-mediated inhibition in the cortex, the network generated a slow oscillation at 2–3 Hz, with field potentials similar to SW (Fig. 5*B*). Field potentials displayed one or several negative/positive sharp deflections, followed by a slowly developing positive wave (Fig. 5*B*, *bottom*). During the spike all cells fired prolonged high-frequency discharges in synchrony, whereas the wave was coincident with neuronal silence in all cell types. This portrait is typical of experimental recordings of cortical and thalamic cells during SW patterns (Pollen, 1964; Steriade, 1974; Avoli et al., 1983; McLachlan et al., 1984; Buzsáki et al., 1990; Inoue et al., 1993). Some TC cells stayed hyperpolarized during the entire oscillation (second TC cell in Fig. 5*B*), as also was observed experimentally (Steriade and Contreras, 1995). A similar oscillation arose if GABA<sub>A</sub> receptors were suppressed in the entire network (data not shown).

These simulations thus indicate that spindles can be transformed into an oscillation with field potentials displaying SW and that this transformation can occur by the alteration of cortical inhibition with no change in the thalamus, in agreement with SW discharges obtained experimentally by diffuse application of diluted penicillin onto the cortex (Gloor et al., 1977). The mechanism of the  $\sim$ 3 Hz oscillation of this model depends on a thalamocortical loop in which both cortex and thalamus are necessary, but none of them generates the 3 Hz rhythmicity alone (see next section below). Figure 3. Corticothalamic feedback can force thalamic circuits into ~3 Hz oscillations because of the properties of GABA<sub>B</sub> receptors. A, Scheme of connectivity and receptor types in a circuit of thalamocortical (TC) and thalamic reticular (RE) neurons. Corticothalamic feedback was simulated through AMPAmediated synaptic inputs (shown on the *left* of the connectivity diagram; total conductance was 1.2  $\mu$ S to RE cells and 0.01  $\mu$ S to TC cells). The *inset* shows simulated burst responses of TC and RE cells after current injection (pulse of 0.3 nA during 10 msec for RE and -0.1 nA during 200 msec for TC). B. A single stimulation of corticothalamic feedback (arrowhead) entrained the circuit into a 10 Hz mode similar to that for spindle oscillations. C, With a strong-intensity stimulation at 3 Hz (arrowheads; 14 spikes per stimulus), RE cells were recruited into large bursts, which evoked IPSPs onto TC cells dominated by GABA<sub>B</sub>-mediated inhibition. In this case the circuit could be entrained into a different oscillatory mode, with all cells firing in synchrony. D, Weak stimulation at 3 Hz (arrowheads) entrained the circuit into spindle oscillations (identical intensity as in B). E, Strong stimulation at 10 Hz (arrowheads) led to quiescent TC cells because of sustained GABA<sub>B</sub> current (identical intensity as in C).



The progressive transformation between spindles and SW oscillations in the model is shown in Figure 6. With intact cortical inhibition the discharge of cells in the network was limited to a few spikes. Consequently, IPSPs in PY cells were almost exclusively GABA<sub>A</sub>-mediated, leading to field potentials consisting of negative deflections only (Fig. 6, 100%). With the intracortical inhibition partially reduced, there was an increased tendency of producing prolonged discharges and an increased contribution of GABA<sub>B</sub> IPSPs in PY cells, leading to small positive waves in field potentials (Fig. 6, 50%). With a further reduction of intracortical GABA<sub>A</sub>-mediated inhibition, the system showed fully developed SW complexes in field potentials, with oscillation frequencies within the 2–3 Hz range (Fig. 6, from 25 to 0%). The frequency of SW oscillations was approximately proportional to the amount of fast inhibition still present in the cortex. The occurrence of a positive spike also was correlated with intracortical fast inhibition (Fig. 6), in agreement with the effect of  $GABA_A$  conductances in Figure 2B.

The waxing and waning appearance of spindles (Fig. 6, 100%) was attributable here to intrathalamic mechanisms. A calciumdependent upregulation of  $I_h$  in TC cells was included here, similar to previous models (Destexhe et al., 1993, 1996a). Such regulation was demonstrated recently in thalamic slices (Lüthi and McCormick, 1998). This mechanism was responsible for the waxing and waning of oscillations in model thalamic and thalamocortical networks (Destexhe et al., 1996a, 1998a). It interesting to note that SW oscillations also may follow a similar waxing and waning envelope (Fig. 6, 25%), which was attributable here to the same intrathalamic mechanisms as spindles. The model therefore suggests that the calcium-dependent upregulation of  $I_h$  in TC cells is responsible for the temporal modulation of SW oscillations and may lead to bursts of several cycles of SW oscillations, interleaved with long periods of silence (~20 sec), as are observed experimentally in sleep spindles and SW epilepsy, thus stressing further the resemblance between the two types of oscillation.

# A thalamocortical loop mechanism for spike-and-wave oscillations

The thalamocortical mechanism leading to SW oscillations in this model is illustrated and compared with spindles in Figure 7. During spindles the oscillation is generated by intrathalamic interactions (TC-RE loop in Fig. 7A). Oscillations can also be generated by a thalamocortical loop (TC-Cx-RE loop in Fig. 7A), as suggested previously (Destexhe et al., 1998a). The combined action of intrathalamic and thalamocortical loops provides a moderate excitation of RE cells, which evokes GABA<sub>A</sub>mediated IPSPs in TC cells and sets the frequency to  $\sim 10$  Hz. During SW oscillations (Fig. 7B) an increased cortical excitability provides a corticothalamic feedback that is strong enough to force prolonged burst discharges in RE cells, which in turn evoke IPSPs in TC cells dominated by the GABA<sub>B</sub> component. In this case the prolonged inhibition sets the frequency to  $\sim$ 3 Hz. The oscillation is generated by a thalamocortical loop (TC-Cx-RE loop in Fig. 7B) in which the thalamus is intact. Therefore, if the cortex is inactivated during SW, this model predicts that the thalamus



Figure 4. Transformation of spindle oscillations into ~4 Hz oscillations by blocking thalamic inhibition in thalamocortical networks. A, Scheme of the connectivity among different cell types: 100 cells of each type were simulated, including thalamocortical (TC) and thalamic reticular (RE) cells, cortical pyramidal cells (PY), and interneurons (IN). The connectivity is shown by continuous arrows, representing AMPA-mediated excitation, and dashed arrows, representing mixed GABA<sub>A</sub> and GABA<sub>B</sub> inhibition. In addition, PY cells were interconnected by using AMPA receptors, and RE cells were interconnected by using GABAA receptors. The inset shows the repetitive firing properties of PY and IN cells that follow depolarizing current injection (0.75 nA during 200 msec; -70 mV rest). B, Spindle oscillations in the thalamocortical network in control conditions. Five cells of each type, equally spaced in the network, are shown (0.5 msec time resolution). The field potentials, consisting of successive negative deflections at  $\sim 10$  Hz, are shown at the bottom. C, Oscillations after the suppression of GABAA-mediated inhibition in thalamic cells with cortical inhibition intact (all GABA<sub>A</sub> conductances postsynaptic to RE cells were suppressed). The network generated synchronized oscillations at  $\sim$ 4 Hz, with thalamic cells displaying prolonged discharges. PY cells showed discharge patterns similar to those of spindles but at a slower frequency; so did the field potentials (bottom).

should resume generating spindle oscillations, as observed experimentally in cats treated with penicillin (Gloor et al., 1979).

The relation between cellular events and field potentials in this model of SW is shown in Figure 8. The pattern displayed by the network is similar to Figure 1*C*2: high-frequency discharges generated spike components in the field potentials, whereas wave components were generated by GABA<sub>B</sub> IPSPs in PY cells because of the prolonged firing of cortical interneurons. The hyperpolarization of PY cells during the wave also contained a significant contribution from the voltage-dependent K<sup>+</sup> current  $I_{\rm M}$  (data not shown), maximally activated because of the prolonged discharge of PY cells during the spike. The wave component is therefore attributable in this model to two types of K<sup>+</sup> currents, intrinsic and GABA<sub>B</sub>-mediated. The relative contribution of each current to the wave depends on its respective conductance values.

During the spike component the discharges were not perfectly in phase. As indicated in Figure 8*B*, there was a significant phase advance of TC cells, as observed experimentally (Inoue et al., 1993). This phase advance was responsible for the initial negative spike in the field potentials, which coincided with the first spike in the TC cells (Fig. 8*B, dashed line*). This feature implements the precedence of EPSPs over IPSPs in the PY cell to generate SW complexes, as evidenced above (see Fig. 2*A*). The simulations therefore suggest that the initial spike of SW complexes is attributable to thalamic EPSPs that precede other synaptic events in PY cells.

### Determinants of spike-and-wave oscillations

The critical factors involved in the genesis of SW oscillations in the thalamocortical model were characterized by investigating the range of synaptic conductances giving rise to SW for each type of connection in the absence of intracortical GABA<sub>A</sub>-mediated inhibition (Table 1). It must be noted that this model considered greatly simplified single-compartment models of thalamic and cortical neurons, with minimal sets of intrinsic currents, no dendrites (and therefore no dendritic currents and no dendritic synapses), and simplified models of intrinsic and synaptic currents. The conductance values therefore cannot match quantitatively the physiological values and must be interpreted qualitatively.

Figure 5. Transformation of spindle oscillations into ~3 Hz oscillations with spikeand-wave field potentials by reducing cortical inhibition. Shown is a similar arrangement of traces as in Figure 4, B and C. A, Oscillations with a 50% decrease of GABA<sub>A</sub>-mediated inhibition in cortical cells (0.075  $\mu$ S, IN $\rightarrow$ PY). Stronger burst discharges appeared within spindle oscillations, leading to large-amplitude negative spikes, followed by small positive waves in the field potentials (bottom). B, Oscillations after suppression of GABAA-mediated inhibition in cortical cells. All cells displayed prolonged discharges in phase, separated by long periods of silences, at a frequency of  $\sim 2$  Hz. GABA<sub>B</sub> currents were activated maximally in TC and PY cells during the periods of silence. Field potentials (bottom) displayed spike-and-wave complexes. Thalamic inhibition was intact in A and B.

Table 1 shows the optimal values of the conductance that were used and, for each connection, the range of values leading to SW oscillations. The minimal frequency of SW bursts when each parameter was varied within 50-200% of the optimal value is indicated in the last two columns of Table 1. The synaptic conductances that were influential on SW were  $PY \rightarrow PY$ ,  $PY \rightarrow IN$ ,  $IN \rightarrow PY, RE \rightarrow RE, RE \rightarrow TC (GABA_B), PY \rightarrow RE, and a weak$ effect for RE $\rightarrow$ TC (GABA<sub>4</sub>). TC $\rightarrow$ RE, TC $\rightarrow$ PY, TC $\rightarrow$ IN, and PY→TC had minimal effect. As expected, the recurrent excitation between pyramidal cells ( $PY \rightarrow PY$ ) and the excitation of interneurons (PY $\rightarrow$ IN), as well as the inhibitory feedback on PY cells (IN $\rightarrow$ PY), are effective on SW because these conductance determine the excitability of the cortical network. Less expected was the role of cortical excitatory feedback on RE cells (PY $\rightarrow$ RE), intra-RE inhibition (RE $\rightarrow$ RE), and the GABA<sub>B</sub> inhibition from RE onto TC cells (RE $\rightarrow$ TC). These factors are examined in more detail below.

A first influential factor was the intra-RE GABAergic connections. Figure 9A shows the transition curve from SW oscillations to spindle waves as a function of intracortical GABA<sub>A</sub> inhibition, similar to Figure 6. Reinforcing intra-RE GABA<sub>A</sub> inhibition significantly reduced SW in favor of the spindles (Fig. 9A, compare *open triangles* with *filled circles*), whereas decreasing this inhibition had the opposite effect (Fig. 9A, *open squares*). In the model, reinforcing intra-RE GABA<sub>A</sub>-mediated inhibition diminished the tendency of RE cells to produce bursts of action potentials, therefore diminishing GABA<sub>B</sub>-mediated IPSPs in TC cells and reducing the tendency to generate SW oscillations. This



behavior is consistent with the presumed role of the anti-absence drug clonazepam, which may reduce the tendency of the network to produce SW by specifically acting on  $GABA_A$  receptors in the thalamic RE nucleus (Huguenard and Prince, 1994a; Gibbs et al., 1996; Hosford et al., 1997).

A second factor that was particularly effective on SW was the corticothalamic feedback on RE cells. Reducing the AMPA conductance of cortical EPSPs on RE cells significantly diminished SW in favor of the spindles (Fig. 9*B*, compare *open triangles* with *filled circles*), and increasing this conductance had the opposite effect. The model therefore indicates that diminishing the impact of corticothalamic EPSPs on RE cells is a potential factor in reducing the threshold for SW in the network.

The need for larger conductances of cortical EPSPs in RE cells versus TC cells also is evidenced for SW oscillations, similar to a previous suggestion in the context of spindle oscillations (Destexhe et al., 1998a). SW oscillations coexisting with spindles required at least four times larger AMPA conductances on RE cells ( $0.4 \,\mu$ S for PY $\rightarrow$ RE and  $0.1 \,\mu$ S of PY $\rightarrow$ TC in Table 1). This is consistent with the anatomical observation that cortical synapses contact only the distal dendrites of TC cells (Liu et al., 1995), leading to attenuated cortical EPSPs.

An additional influential factor, not included in Table 1, was the T-current conductance in RE cells. Reducing the T-current of RE cells significantly reduced SW in favor of the spindles (Fig. 9C, compare open triangles with filled circles), whereas reinforcing this current had the opposite effect (Fig. 9C, open squares). Reducing T-current amplitude therefore diminishes the tendency of the



Figure 6. Gradual transformation of spindles to spike-and-wave complexes. The field potentials obtained for different simulations similar to Figure 5 are shown from *top* to *bottom*. The different simulations correspond to identical conditions, except that intracortical GABA<sub>A</sub>-mediated inhibition (IN $\rightarrow$ PY) was reduced, with total conductance values of 0.15  $\mu$ S (100%), 0.075  $\mu$ S (50%), 0.0375  $\mu$ S (25%), 0.018  $\mu$ S (12%), and 0.009  $\mu$ S (6%). 100% corresponded to a spindle sequence (same simulation as in Fig. 4*B*) and 0% to fully developed SW complexes when the intracortical GABA<sub>A</sub> inhibition was suppressed (same simulation as in Fig. 5*B*); intrathalamic inhibition was intact in all cases.

network to produce SW, similar to reinforcing GABAergic inhibition in the RE nucleus. This effect is consistent with the experimental finding that the T-current is increased selectively in RE cells in a rat model of absence epilepsy (Tsakiridou et al., 1995).

On the other hand, reducing the T-current conductance in TC cells had only a weak effect on SW threshold (data not shown), but T-current reduction >40% in TC cells led to the suppression of oscillatory behavior. This was consistent with the effect of the anti-absence drug ethosuximide in reducing the total T-current conductance in TC cells (Coulter et al., 1989).

As predicted from the mechanism of Figure 7*B*, the frequency of SW essentially was determined by GABA<sub>B</sub>-mediated IPSPs on TC cells (Fig. 9*D*, *filled circles*). Changing the decay of intrathalamic GABA<sub>B</sub> currents (parameter  $K_4$ ) affected only the frequency, with minimal changes in the bursting patterns of the different cell types (data not shown). This effect was attributable to the fact that, in this model, the duration of the wave is determined essentially by GABA<sub>B</sub> IPSPs in TC cells, longer IPSPs leading to slower SW by further delaying the rebound of



Figure 7. Thalamocortical loop mechanism for spike and wave. Simplified diagrams represent the major steps involved in generating oscillations (*Cx*, cortex). *A*, Spindle oscillations resulting from a mutual recruitment of thalamic TC and RE cells (*thick lines*) in which TC cells rebound after fast GABA<sub>A</sub>-mediated IPSPs, setting the frequency to ~10 Hz. Here, the oscillation is generated in the thalamus and is reinforced by the thalamocortical loop (*thin lines*). *B*, Proposed mechanism for spike and wave. In this case the corticothalamic feedback is much stronger because of increased cortical excitability, forcing thalamic cells to display prolonged burst discharges, which evoke GABA<sub>B</sub>-mediated IPSPs in TC cells. This prolonged inhibition prevents cells from firing during ~300 msec and sets the frequency to ~3 Hz.

TC cells. The frequency varied from 1 to 5 Hz for decay values of 50-250% of the control value, suggesting that the different frequency of SW bursts in different experimental models may be attributable to differences in the kinetics of GABA<sub>B</sub>-mediated inhibition in TC cells.

The T-current amplitude in TC cells also affected the SW frequency (Fig. 9D, open squares). Stronger T-current conductances led to earlier rebound and faster frequencies. By contrast, the T-current amplitude in RE cells had minimal effect on SW frequency (Fig. 9C, open triangles). Consistent with the mechanism depicted in Figure 7B, the frequency of SW was mostly attributable to intrathalamic mechanisms, whereas the threshold for SW was dependent on the different elements involved in the thalamus–cortex–thalamus loop.

### DISCUSSION

This paper proposed a thalamocortical loop mechanism for the genesis of spike-and-wave oscillations. This mechanism, its simi-



*Figure 8.* Phase relations during simulated spike-and-wave discharges. *A*, Local field potentials (*LFP*) and representative cells of each type during SW oscillations. *Spike*, All cells displayed prolonged discharges in synchrony, leading to spiky field potentials. *Wave*, The prolonged discharge of *RE* and *IN* neurons evoked maximal GABA<sub>B</sub>-mediated IPSPs in *TC* and *PY* cells, respectively (*dashed arrows*), stopping the firing of all neuron types during a period of 300–500 msec and generating a slow positive wave in the field potentials. The next cycle restarted because of the rebound of *TC* cells after the GABA<sub>B</sub> IPSP (*arrow*). *B*, Phase relationships in the thalamocortical model. *TC* cells discharge first, followed by *PY*, *RE*, and *IN* cells. The initial negative peak in the field potentials coincided with the first spike in *TC* cells before the *PY* cells started firing and was generated by thalamic EPSPs in *PY* cells.

larities and differences with experimental SW, plus predictions to test its validity are discussed successively.

### A GABA<sub>B</sub>-based mechanism for spike and wave

The cellular mechanism proposed here is based on the following properties:

(1) Because of the characteristics of  $GABA_B$ -mediated responses, simulated field potentials can display SW waveforms if cortical pyramidal cells and interneurons generate prolonged discharges in synchrony, without the need of any other assumption about intrinsic cellular or circuit mechanisms.

(2) Also because of the characteristics of  $GABA_B$ -mediated inhibition, model thalamic circuits can be forced into ~3 Hz oscillations. It is known from slice experiments that thalamic circuits naturally oscillate at ~10 Hz but display ~3 Hz oscillations in the presence of  $GABA_A$ -receptor agonists (von Krosigk et al., 1993). The present model suggests that a similar oscillation can be forced in *intact* thalamic circuits if corticothalamic feedback EPSPs are strong enough.

(3) Generalized  $\sim$ 3 Hz oscillations can be generated through thalamocortical loops. If because of an increase of cortical excitability the thalamic-projecting cortical cells generate exceedingly strong discharges, then the ensuing corticothalamic feedback EPSPs may become strong enough to force the thalamus in the 3

Hz mode. The  $\sim$ 3 Hz oscillations then invade the entire network through thalamocortical loops. The  $\sim$ 3 Hz frequency depends on intrathalamic GABA<sub>B</sub>-mediated inhibition.

(4) This  $\sim$ 3 Hz oscillation generates SW field potentials. During the spike the thalamic and cortical cells produce prolonged discharges in synchrony, whereas the wave is generated by a mixture of voltage-dependent and GABA<sub>B</sub>-mediated K<sup>+</sup> currents.

# Similarities with experimental models of spike and wave

This thalamocortical loop model is consistent with a number of experimental results on SW epilepsy: (1) thalamic and cortical neurons discharge in synchrony during the spike, whereas the wave is characterized by neuronal silence (Pollen, 1964; Steriade, 1974; Avoli et al., 1983; McLachlan et al., 1984; Buzsáki et al., 1990; Inoue et al., 1993), similar to the data in Figures 4*E* and 8*A*; (2) TC cell firing precedes that of other cell types, followed by cortical cells and RE cells (Inoue et al., 1993), similar to the phase relations of the present model (see Fig. 8B); (3) SW patterns disappear after the removal of either the cortex (Avoli and Gloor, 1982) or the thalamus (Pellegrini et al., 1979; Vergnes and Marescaux, 1992), as also predicted by the present mechanism; (4) antagonizing thalamic GABA<sub>B</sub> receptors suppresses SW discharges (Liu et al., 1992), consistent with this model; (5) spindle oscillations can be transformed gradually into SW discharges (Kostopoulos et al., 1981a,b), as described in Figure 6.

The present mechanism also emphasizes a critical role for the RE nucleus. Reinforcing  $GABA_A$ -mediated inhibition in the RE nucleus will antagonize the genesis of large burst discharges in RE cells by corticothalamic EPSPs, antagonizing the genesis of  $GABA_B$ -mediated IPSPs in TC cells and therefore antagonizing SW. This property is consistent with the diminished frequency of seizures that is observed after the reinforcement of  $GABA_A$  receptors in the RE nucleus (Liu et al., 1991). It is also consistent with the action of the anti-absence drug clonazepam, which seems to act preferentially by enhancing  $GABA_A$  responses in the RE nucleus (Hosford et al., 1997), leading to diminished  $GABA_B$ -mediated IPSPs in TC cells (Huguenard and Prince, 1994a; Gibbs et al., 1996).

The fact that injections of GABA<sub>A</sub> antagonists in the thalamus with intact cortex failed to generate SW (Ralston and Ajmone-Marsan, 1956; Gloor et al., 1977; Steriade and Contreras, 1998) also was considered. In the model, suppressing thalamic GABA<sub>A</sub> receptors led to "slow spindles" at ~4 Hz, very different from SW oscillations (see Fig. 4*C*). In this case the discharge of PY cells was extremely brief, because cortical GABA<sub>A</sub>-mediated inhibition was preserved and no GABA<sub>B</sub> IPSPs could be evoked. This result is consistent with the powerful control exerted on pyramidal cells by intracortical GABA<sub>A</sub>-mediated inhibition, as shown by intracellular recordings and modeling (Contreras et al., 1997).

### Differences with experimental models of spike and wave

On the other hand, a number of experimental observations are not consistent with the mechanism presented here. First, an apparent intact cortical inhibition was reported in cats treated with penicillin (Kostopoulos et al., 1983). However, this study did not distinguish between GABA<sub>A</sub> and GABA<sub>B</sub>-mediated inhibition. In the present model, even when GABA<sub>A</sub> was antagonized, IPSPs remained approximately the same size because cortical interneurons fired stronger discharges (see Fig. 4D,E) and led to stronger GABA<sub>B</sub>- and GABA<sub>B</sub>- and effect between GABA<sub>A</sub> and compensation effect between GABA<sub>A</sub> and

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Receptor type	Location	Optimal value	SW range	SW frequency (50%) in Hz	SW frequency (200%) in Hz
AMPA	PY→PY	0.6 µS	0.3–0.9 µS	3.0	_
AMPA	PY→IN	$0.2 \mu S$	0.06–2 µS	1.3	1.9
$GABA_{R}$	IN→PY	0.03 µS	0.02–5 µS*	1.3	2.0
AMPA	TC→RE	$0.2 \mu S$	$0-5 \mu S^*$	1.7	1.7
$GABA_A$	RE→RE	0.2 µS	0–1.2 μS	1.4	2.5
GABA <sub>4</sub>	RE→TC	0.02 µS	0–1 µS	1.7	1.8
GABA <sub>B</sub>	RE→TC	0.04 µS	0.01–5 µS*	2.1	1.3
AMPA	TC→PY	1.2 μS	0.15–5 μS*	1.7	1.7
AMPA	TC→IN	$0.4 \mu S$	$0-5 \mu S^*$	1.7	1.7
AMPA	PY→RE	$1.2 \mu S$	0.4–5 µS*	2.2	1.5
AMPA	PY→TC	0.01 µS	0-0.1 µS	1.7	1.7

Conductance values represent the sum of all individual synaptic conductances of the same type converging to a given cell. The range of conductance values giving rise to SW oscillations is indicated in the fourth column (\*no value >5  $\mu$ S was tested, because sodium spike inactivation may occur for too high conductance values). The minimal oscillation frequency is indicated in the last two columns when each respective conductance was set to 50 and 200% of the optimal value; a value equal to control (1.7 Hz) indicates that this parameter had no detectable influence on SW. All simulations were run by using the thalamocortical network model with suppressed GABA<sub>A</sub>-mediated inhibition in cortical cells (same conditions as in Fig. 5B).



Figure 9. Determinants of spike-and-wave oscillations. A, Effect of GABA<sub>A</sub>-mediated inhibition between RE cells. The lowest frequency of SW complexes is represented as a function of the amount of GABAA inhibition in cortex (simulations similar to Fig. 6). In control (filled circles) the frequency of SW increased steadily up to 60% of cortical GABA<sub>A</sub>; then a transition occurred to spindle oscillations (lowest frequency of  $\sim 8$  Hz). With twice smaller intra-RE GABA<sub>A</sub> conductances (open squares) this transition occurred at  $\sim$ 75% cortical GABA<sub>A</sub>. When intra-RE GABAA conductances were doubled, the domain of SW was significantly smaller, with a transition occurring at  $\sim 20\%$  of cortical GABA<sub>A</sub> (open triangles). B, Effect of corticothalamic feedback on RE cells. With diminished AMPA conductance in PY→RE synapses (50% of control value), the domain of SW was reduced significantly (open triangles), whereas reinforced cortical EPSPs had the opposite effect (open squares). Filled circles, Same control as in A. C, Effect of the T-current conductance in RE cells. With reinforced T-current (200% of control value) the transition occurred at  $\sim$ 75% of cortical GABA<sub>A</sub> (open squares), whereas with diminished T-current (50% of control value) the domain of SW was reduced significantly (open triangles). Filled circles, Same control as in A. D, Determinants of SW frequency. The frequency of SW bursts in the simulation of Figure 5B was represented when several parameters were varied. These parameters are represented as the

percentage of their control value (100% = control). The parameters represented are the decay of intrathalamic GABA<sub>B</sub> currents (*filled circles*), the T-current conductance in TC (*open squares*), and RE cells (*open triangles*).

GABA<sub>B</sub>-mediated IPSPs (data not shown), which may lead to the misleading observation that inhibition is preserved.

Second, some GABA<sub>A</sub> agonists, like barbiturates, may increase the frequency of seizures (Vergnes et al., 1984), possibly via interactions with GABA<sub>A</sub> receptors in TC cells (Hosford et al., 1997). A similar effect was seen in the model (Table 1, GABA<sub>A</sub> RE $\rightarrow$ TC), but this effect was weak. To be simulated more precisely, this type of data would require modeling the variants of GABA<sub>A</sub> receptor types in different cells and addressing how the threshold for SW discharges is affected by various types of GABAergic conductances. These points should be considered in future models. Third, the present model investigated only a thalamocortical loop scenario for the genesis of SW oscillations, but other mechanisms are possible. Although most experimental data are in favor of a mechanism involving both thalamus and cortex (see introductory remarks), numerous experimental evidence also points to a possible intracortical mechanism for SW. Experiments revealed SW in isolated cortex or athalamic preparations in cats (Marcus and Watson, 1966; Pellegrini et al., 1979; Steriade and Contreras, 1998). However, this type of paroxysmal oscillation had a different morphology and was slower in frequency compared with the typical "thalamocortical" SW (1–2.5 vs 3.5–5 Hz; Pellegrini et al., 1979). By contrast, intracortical SW was not

observed in athalamic rats (Vergnes and Marescaux, 1992). Because no intracellular recordings were made during the presumed SW in cat isolated cortex, it is not clear if this oscillation represents the same SW paroxysm as in the intact thalamocortical system. Nevertheless, the cortex is known to display intrinsic oscillations generated by bursting neurons (Silva et al., 1991) and also contains rebound-bursting pyramidal cells in some cortical areas (de la Pena and Geijo-Barrientos, 1996). It may be that these properties are sufficient to sustain a form of purely cortical SW, via a sequence of GABA<sub>B</sub> IPSPs and rebound, similar to the mechanism analyzed here. When more precise experimental data become available, such as intracellular recordings, possible intracortical mechanisms for SW should be investigated by future models.

#### Predictions

Several predictions are generated by this model. First, the wave component of SW was generated by massive  $K^+$  currents, mostly because of GABA<sub>B</sub> receptor activation. This could be observable by performing intracellular recordings during SW while blocking GABA<sub>B</sub> responses. At the network level the injection of GABA<sub>B</sub> antagonists or  $K^+$  currents blockers on the cortex should lead to significant alteration of the wave component in field potentials. One must, however, bear in mind that other mechanisms not taken into account here also may participate to the wave component, such as the activation of Ca<sup>2+</sup> spikes and Ca<sup>2+</sup>-dependent K<sup>+</sup> currents, possibly in dendrites (Traub and Miles, 1991).

A second prediction is that intact thalamic circuits can be forced into a  $\sim$ 3 Hz oscillation by strong stimulation of corticothalamic feedback fibers. This experiment could be performed in slices or in decorticated animals by stimulating the internal capsule. The intensity should be high enough to force large bursts in RE cells, evoking GABA<sub>B</sub> IPSPs in TC cells and delaying their rebound by  $\sim$ 300 msec. Two stimulation patterns are possible: either the stimulation period should match the delay to rebound, or the stimulation could be triggered by the multiunit discharge of TC cells. In the latter case the model predicts a switch from  $\sim$ 10 Hz oscillations to  $\sim$ 3 Hz when the stimulation intensity is increased.

Finally, the model predicts a critical role for the dependence of  $GABA_B$  IPSPs on the number of presynaptic spikes. This effect was simulated by assuming that the binding of four G-proteins is required to activate the K<sup>+</sup> channels underlying GABA<sub>B</sub> responses (Destexhe and Sejnowski, 1995). In the thalamocortical model the dependence on the number of spikes is critical for generating the ~3 Hz oscillations as well as the spike-and-wave waveform in field potentials. Dual intracellular recordings between cortical inhibitory interneurons and their targets should shed light into this question in the near future (A.M. Thomson and A. Destexhe, unpublished data).

In conclusion, this paper suggests a thalamocortical loop mechanism for spike and wave, based on the intrinsic and synaptic properties of thalamic and cortical cells, the characteristics of which are consistent with several experimental models of SW as well as with thalamic slice experiments. The key biophysical components of this mechanism are the activation properties of GABA<sub>B</sub> receptors, combined with the complex intrinsic firing properties of thalamic cells. The model also emphasizes a major role for corticothalamic feedback in triggering powerful bursts in RE cells, by which the cortex can force the thalamus to generate oscillations at  $\sim$ 3 Hz by activating intrathalamic GABA<sub>B</sub>-mediated inhibition. Because thalamic RE cells may generate bursts through dendritic T-currents (Destexhe et al., 1996b), their sensitivity to corticothalamic feedback EPSPs therefore may be maximal (Contreras et al., 1993), leading to the prediction that this structure should be a major target for a possible suppression of seizures.

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