Signal Transfer in Passive Dendrites with Nonuniform Membrane Conductance

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In recent years it became clear that dendrites possess a host of ion channels that may be distributed nonuniformly over their membrane surface. In cortical pyramids, for example, it was demonstrated that the resting membrane conductance $G_m(x)$ is higher (the membrane is “leakier”) at distal dendritic regions than at more proximal sites. How does this spatial nonuniformity in $G_m(x)$ affect the input-output function of the neuron? The present study aims at providing basic insights into this question. To this end, we have analytically studied the fundamental effects of membrane non-uniformity in passive cable structures.

Keeping the total membrane conductance over a given modeled structure fixed (i.e., a constant number of passive ion channels), the classical case of cables with uniform membrane conductance is contrasted with various nonuniform cases with the following general conclusions. (1) For cylindrical cables with “sealed ends,” monotonic increase in $G_m(x)$ improves voltage transfer from the input location to the soma. The steeper the $G_m(x)$, the larger the improvement. (2) This effect is further enhanced when the stimulation is distal and consists of a synaptic input rather than a current source. (3) Any nonuniformity in $G_m(x)$ decreases the electrotonic length, $L$, of the cylinder. (4) The system time constant $\tau$ is larger in the nonuniform case than in the corresponding uniform case. (5) When voltage transients relax with $\tau$, the dendritic tree is not isopotential in the nonuniform case, at variance with the uniform case. The effect of membrane nonuniformity on signal transfer in reconstructed dendritic trees and on the $I/f$ relation of the neuron is also considered, and experimental methods for assessing membrane nonuniformity in dendrites are discussed.

Key words: cable theory; nonuniform membrane conductances; dendritic ion channels; dendritic signal transfer; compartmental modeling; dendritic transients

Transmembrane ion channels are the main carriers of electrical currents in neurons. Their type, kinetics, and spatial distribution may critically determine the properties of the electrical signals that are initiated and propagated in neurons. Importantly, these ion channels are distributed nonuniformly over the neuron surface. An obvious example is the myelinated axons where Ranvier nodes bear a high density of $\mathrm{Na}^+$ channels, whereas the internodes are bare of these channels (Hille, 1992). This spatial nonuniformity has functional implications for the propagation speed of the action potential along the axon. Ion channels are distributed nonuniformly also over the dendritic membrane. Are there some design principles that govern the distribution of channels and optimize certain aspects of signal processing in dendrites?

Early studies on excitatory properties of dendrites can be found in Lorente de Nó (1947), Spencer and Kandel (1961), Llinás and Sugimori (1980), and Schwindt and Crill (1995); for review, see Mel (1994). Recent experimental studies using infrared DIC video microscopy, combined with patch-clamp techniques (Stuart et al., 1993; Dodt and Zieglgansberger, 1994) directly showed that, indeed, ion channels were distributed nonuniformly over the dendritic surface (for review, see Johnston et al., 1996). Recordings from membrane patches, excised from the apical trunk of hippocampal CA1 pyramidal neurons, showed that both the A-type and the delayed rectifier $K^+$ conductances linearly increased with the distance from the soma (Hoffman et al., 1997). The density of $I_h$ currents also increase in these cells (Magee, 1998). An increase in the density of both $I_h$ current and the cesium-insensitive membrane conductance at distal dendritic regions was also recently observed in neocortical layer V pyramids (Stuart and Spruston, 1998). Nonuniformity in dendritic excitability was also found in other neuron types (Stuart and Häusser, 1994; Turrigiano et al., 1995; Bischofberger and Jonas, 1997; Kavalali et al., 1997; for review, see Segev and Rall, 1998).

Spatial distribution and spatiotemporal activation pattern of transmitter-gated conductances may also cause nonuniformity of the dendritic membrane. Thus, the dendritic membrane conductance is expected to develop time- and activity-dependent spatial heterogeneities (Holmes and Woody, 1989).

Most of our intuitions regarding how the interplay between dendritic morphology, physiology, and input conditions determine the input–output properties of dendrites rely on Rall’s cable theory. In this framework, the effect of geometric nonuniformities in the dendritic tree, including tapering and branching, was analyzed (Butz and Cowan, 1974; Horwitz, 1981, 1983; Poznanski,
Table 1. Table of symbols

| Symbol | Definition |
|--------|------------|
| $V$    | Transmembrane voltage (relative to resting potential) | mV |
| $d$    | Diameter of the dendritic cylinder | cm |
| $\ell$ | Length of the dendritic cylinder | cm |
| $x$    | Physical distance from the origin (usually the soma) | cm |
| $G_m(x)$ | Specific membrane conductance | S/cm² |
| $\bar{G}_m = 1/\ell \int_0^\ell G_m(x)dx$ | Average membrane conductance | S/cm² |
| $\gamma_m(x) = G_m(x)/\bar{G}_m$ | Dimensionless membrane conductance | Dimensionless |
| $R_m(x)$ | Specific membrane resistance | $\Omega$ cm² |
| $R_i$  | Specific cytoplasm resistivity | $\Omega$ cm |
| $\tau_i = 4R_i/(md^2)$ | Internal resistance per unit length of cylinder | $\Omega$ cm |
| $C_m$  | Specific membrane capacitance | $F$/cm² |
| $\alpha$ | Slope parameter ($-1 \leq \alpha \leq 1$) | Dimensionless |
| $\lambda(x) = \sqrt{d/4G_m(x)\bar{R}_i}$ | Space constant (nonuniform case) | cm |
| $\lambda^u = \sqrt{d/4G_m^u\bar{R}_i}$ | Space constant (uniform case) | cm |
| $L = \int_0^\ell dx/\lambda(x)$ | Generalized electrotonic length | Dimensionless |
| $L^u = \ell/\lambda^u$ | Electrotonic length (uniform case) | Dimensionless |
| $X^u = x/\lambda^u$ | Distance from location $x$ to origin, in units of the uniform space constant | Dimensionless |
| $R_m(x)$ | Input resistance at location $x$ | $\Omega$ |
| $G_m(x) = 1/R_m(x)$ | Input conductance at location $x$ | $S$ |
| $R_{xy}$ | Transfer resistance from location $x$ to location $y$ | $\Omega$ |
| $I_x$ | Axial current | $A$ |
| $G_x = (\pi/2) \sqrt{4G_m(x)/\bar{R}_i}$ | Input conductance of a semi-infinite uniform cylinder | $S$ |
| $G_r$ | Axial leak conductance at boundary | $S$ |
| $B = G_x/G_r$ | Boundary condition | Dimensionless |
| $\tau_m(x) = C_m/G_m(x)$ | Local membrane time constant | sec |
| $\tau^u_m = C_m/G_m^u$ | Membrane time constant (uniform case) | sec |
| $\tau_0^u, \tau^u_n$ | $n$-th equalizing time constant (uniform and slope case, respectively) | sec |
| $\delta(x, \delta(t))$ | Delta function | |
| $\delta(x,y,t)$ | Green function | |
| $\delta_{syn}$ | Synaptic conductance | S |
| $E_{syn}$ | Synaptic reversal potential | mV |

1988; Schierwagen, 1989; Abbott, 1992; Holmes et al., 1992; Holmes and Rall, 1992a,b; Agmon-Snir and Segev, 1993; Major et al., 1993a–c; Segev et al., 1995; Rall and Agmon-Snir, 1998), and the impact of nonlinear voltage- and transmitter-gated dendritic ion channels, as well as of massive synap tic input on the input-output properties of neurons, was explored theoretically (Barrett and Crill, 1974; Shepherd et al., 1985; Miller et al., 1985; Fleshman et al., 1988; Segev and Rall, 1988; Clements and Redman, 1989; Holmes, 1989; Bernander et al., 1991; Rapp et al., 1992; Siegel et al., 1994; Bernard et al., 1994; De Schutter and Bower, 1994; Pinsky and Rinzel, 1994; Schwindt and Crill, 1995; Stuart and Sakmann, 1995; Wilson, 1995; Mainen and Sejnowski, 1996). A review of these issues can be found in Koch (1999). Experimental studies on the input–output properties of nonlinear dendrites can be found in Laurent et al. (1993), Nicoll et al. (1993), Haag and Borst (1996), and Chen et al. (1997). However, there is yet no systematic study on the effect of membrane nonuniformity in dendrites, although previous studies did consider the possibility that the soma membrane is leakier than the dendritic membrane (Rall, 1962; Rall and Shepherd, 1968; Iansek and Redman, 1973; Durand, 1984; Kawato, 1984; Major et al., 1993a).

Motivated by the recent experimental evidence about membrane nonuniformity in dendrites, the present study explores, using analytical tools, whether there is some functional advantage of membrane nonuniformity for the transfer of synaptic signals from the dendrites to the soma. In the tradition of W. Rall, we treat here, as a first stage, the case in which the dendritic membrane conductance $G_m(x)$ is spatially nonuniform and passive. All mathematical considerations are grouped in the Appendixes (online at www.jneurosci.org).

**MATERIALS AND METHODS**

The cable equation (Rall, 1959) away from current sources for a passive cylinder with nonuniform membrane conductance is:

$$\lambda(x) \frac{\partial^2 V}{\partial x^2} - \tau_m(x) \frac{\partial V}{\partial t} = 0, \quad (1)$$

where $\tau_m(x) = C_m/G_m(x)$ is the (variable) passive membrane time constant and $\lambda(x) = \sqrt{d/4G_m(x)\bar{R}_i}$ is the (variable) space constant, $G_m(x)$ is the specific membrane conductance, and $d$ is the diameter of the cylinder. In the uniform case, in which $G_m(x)$ is constant, $\lambda(x) = \lambda^u$, and $\tau_m(x) = \tau^u_m$; the superscript, “$u$” is used throughout this work to label the uniform case. These and other quantities are defined in Table 1.

To understand the effects of different $G_m(x)$ functions, we need to impose some constraint that will allow us to compare these $G_m(x)$ functions. The most natural assumption is that the total membrane conductance of the cable is preserved in all cases:

$$\pi d \int_0^\ell G_m(x)dx = \text{const.}. \quad (2)$$

This implies that there is a fixed number of (passive) ion channels in the modeled structure and that these channels are allocated to different regions of the cable, depending on the shape of $G_m(x)$.
The case of membrane nonuniformity that will gain the main focus of the present work is the slope case, in which \( G_m(x) \) linearly increases with the distance from the “soma” \((x = 0)\), namely \( G_m(x) = ax + b \). This case has received some recent experimental support (Hoffman et al., 1997). All quantities are labeled, in this case, with the superscript “s.” In the case of cylinder of length \( \ell \), the constraint of a fixed total conductance, Equation 2, implies that:

\[
G_m(x) = G_m \left(1 + 2a \frac{x - \ell/2}{\ell}\right),
\]

where \( G_m \) is the average membrane conductance over the membrane area, \( x \) is the location along the cylinder \((0 \leq x \leq \ell)\), and the dimensionless parameter \( \alpha = a/(2G_m) \) is the slope \( a \) in units of \( \ell G_m \). Because \( G_m(x) \) should be positive for all \( x \), \( a \) is bounded, \(-1 \leq \alpha \leq 1\). The uniform case is obtained when \( \alpha = 0 \), whereas for the case of maximal slope, \( \alpha = 1 \). In this case, the membrane conductance increases from \( 0 \) at \( x = 0 \) to a maximum of \( 2G_m \) at \( x = \ell \).

For DC current injection, the membrane voltage (away from current sources) satisfies the steady-state cable equation:

\[
\lambda^2(x) \frac{d^2V}{dx^2} - V = 0.
\]

Equation 4 can be analytically solved for several nonuniform conductance profiles. The solutions for typical increasing of \( G_m(x) \) functions (power law, exponential, etc.) are given in Appendix B; they all involve special functions (Abramowitz and Stegun, 1970). Mathematica software (Wolfram, 1991) was used for numerical evaluations of these analytical solutions; transient simulations and computations of transfer resistances in reconstructed dendrites were implemented using NEURON simulator (Hines and Carnevale, 1997).

The transient voltage–response of a cable with nonuniform membrane conductance can be computed using the Green function, \( \delta_i(x, y, t) \), of the problem which satisfies:

\[
\left(\lambda^2 \frac{d^2}{dx^2} - \delta \right) \delta_i(x, y, t) = -\delta(x-y)\delta(t),
\]

together with the imposed boundary and initial conditions. Separating time and space variables, the Green function can be expressed as an orthonormal expansion (Rall, 1969):

\[
\delta_i(x, y, t) = \sum_{j=0}^{\infty} c_j(x)\delta_j(y)e^{-\gamma_j t},
\]

where the basis functions \( \delta_j \) satisfy:

\[
\int_0^\ell \delta_j(x)\delta_k(x)dx = \begin{cases} 1 & j = k \\ 0 & j \neq k \end{cases}.
\]

The system time constant \( \tau_0 \) and the equalizing time constants \( \tau_j (j \geq 1) \), together with the associated eigenfunctions \( \delta_j (j \geq 0) \), are obtained by solving the eigenvalue problem:

\[
\lambda^2 \frac{d^2\delta_j}{dx^2} - \frac{G_m(x)}{G_m} \tau_j \delta_j = 0,
\]

with imposed boundary conditions at \( x = 0 \) and \( x = \ell \). It is meaningful to distinguish \( \tau_0 \) from the other time constants (Rall, 1969). Indeed, \( \tau_0 \) governs the slow relaxation of the voltage at large \( t \), and for uniform cables with sealed ends, \( \tau_0 = \tau_d \). The other, smaller time constants control the faster spatial equalization of voltage gradients, and they vanish in the limit of extremely compact cables \((L \to 0)\). As shown in Appendix E, the eigenvalue problem is analytically tractable in the slope case: the eigenvalues are determined by solving a transcendental equation, and the associated eigenfunctions can be expressed as linear combinations of Airy functions.

The eigenvalue problem computed over the slope case is formally identical to the Schrödinger eigenvalue problem for the one-dimensional bounded motion of a particle in quantum mechanics (Landau and Lifschitz, 1997). This analogy, which is elaborated on in Appendix D.1, is useful because it provides a visual representation of the Green function of the cable problem in terms of discrete energy states in a potential well. It provides new insight on how the system time constant and the equalizing time constants, together with the associated eigenfunctions, depend on the membrane conductance profile and on other cable parameters such as the electrotonic length. Note also that another representation of the Green function, as a diffusion process, was proposed by Abbott et al. (1991), Abbott (1992), and Bressloff and Taylor (1993); this approach also relies on concepts and methods (i.e., path integrals or their discrete counterpart) that were previously introduced in physics. These methods can be applied to any profile of nonuniform specific conductance on the dendritic tree, but they require the numerical evaluation of the contributions of many paths. This limits its practical usefulness mostly to the accurate estimation of the short-term transient behavior. In contrast, our approach is limited to certain types of conductance profiles, but it provides analytical results and enables us to accurately determine, through the computation of the larger time constants, how the slow relaxation of transients proceeds.

**RESULTS**

**Steady current input in nonuniform cylinders**

The steady-state voltage profile along a cylinder with both ends sealed, following constant current injection at different locations, is shown in Figure 1A for the uniform case and the maximal slope case \((\alpha = 1)\). This figure shows that the voltage at \( x = 0 \) (soma) is always larger in the slope case, even when the local voltage response at the input site is smaller than in the uniform case (e.g., at the distal end where \( x = \ell = 1000 \mu m \)). As demonstrated in Figure 1B, the voltage attenuation between any input point \( x \) and the soma is smaller in the slope case. Note that this figure also describes the attenuation of the time integral of the potential in the case of transient current injection (Rall and Rinzel, 1973). Thus, distributing the same total membrane conductance nonuniformly rather than uniformly improves the voltage response at the soma.

To characterize the effect of steady current injection at point \( x \) on the voltage response at the soma, the transfer resistance \( R_{\text{trans}}(x) = I(x)/V(x) \) is used. Figure 2A shows the normalized transfer resistance between the input location and the soma for cylinders of different electrotonic length. In the maximal slope case, the transfer resistance is always larger (for every \( x \)) than in the corresponding uniform case, and thus for all input locations the voltage response at the soma is larger when the conductance linearly increases in the cylinder. This property is independent of the cylinder physical length. The difference between the two cases grows with the electrotonic length of the cylinder. It is also true for all the intermediate linear slopes (i.e., \( 0 < \alpha < 1 \)).

Figure 2B shows the normalized input resistance \( R_{\text{in}}(x) = R_{\text{trans}}(x) \) for cylinders with electrotonic length \( L = 1 \). In the uniform case, \( R_{\text{in}}(x) \) is symmetric with respect to the midpoint \( x = \ell/2 \), where the minimum is obtained. This is no longer true in the maximal slope case in which \( R_{\text{in}}(x) \) is larger in the proximal part of the cylinder and smaller in its distal part compared with the uniform case. This is expected because, in the slope case, the membrane is “leakier” in distal regions of the cylinder. Because \( R_{\text{in}} \) is determined not only by the local membrane conductance but also by the cable properties and the boundary conditions, the intersection between \( R_{\text{in}}(x) \) in the two cases is not at the cylinder’s midpoint, \( x = 0.5 \) (where \( G_m(x) = G_m \)), but rather at \( x^* = 0.57 \). A given current input at \( x = 0.57 \) will result in a smaller local voltage response in the slope case because of the smaller input resistance in these sites compared with that of the uniform case. Nevertheless, the resultant voltage at the soma is larger in the slope case because of the larger transfer resistance in this case. In contrast, for all \( x^* < 0.57 \), both \( R_{\text{in}} \) and \( R_{\text{trans}} \) are larger in the slope case, giving rise to a larger voltage response both locally and at the soma. The “benefit,” in terms of the relative increase of the soma...
when increases with the steepness of the conductance profile. Indeed, conductance, and the difference compared with the uniform case transfer resistance is always increased by a gradient of membrane 0. Figure 3 shows that, compared with the uniform case, the conductance (Eq. 2) holds, and in all nonuniform cases, malized such that the constraint of a fixed average membrane 82\% for proximal inputs in cylinders with electrotonic length, than a uniform conductance ranges from 3\% for distal inputs to 26\% at more proximal sites, with an average of 16% for proximal inputs in cylinders with electrotonic length, $L^u = 1$ (see Fig. 4B, current input).

The focus in the above analysis was on the slope case, in which $G_m(x)$ is a linear function of $x$, and more specifically on the limiting case where $\alpha = 1$. In Figure 3, the transfer resistance for several power law functions of $G_m(x)$ with increasing exponent (0, 1/2, 1, and 2) is computed. All cases with integer power are actually analytically tractable (see Appendix B.2). $G_m(x)$ is normalized such that the constraint of a fixed average membrane conductance (Eq. 2) holds, and in all nonuniform cases, $G_m(0) = 0$. Figure 3 shows that, compared with the uniform case, the transfer resistance is always increased by a gradient of membrane conductance, and the difference compared with the uniform case increases with the steepness of the conductance profile. Indeed, when $G_m(x)$ increases with an exponent of 2, the benefit, in terms of the relative increase of the soma voltage, ranges from 6% for distal inputs to 26% at more proximal sites, with an average of 17%. Note also that because of the reciprocity property of the transfer resistance [R$_{s,x} = R_{x,s}$ (Koch, 1999)], the effect of spatially decreasing conductance profiles can be deduced from these results (see Appendix C.3).

Finally, we note that the generalized electrotonic length of a given cylinder, $L = \int_0^1 dx/x$, is smaller for the slope case compared with the uniform case. Indeed, for the slope case, $L$ is given by:

$$L = L^u \frac{(1 + |\alpha|)^{3/2} - (1 - |\alpha|)^{3/2}}{3|\alpha|}.$$  

$L$ decreases with increasing $|\alpha|$ and reaches its minimum, $L = \sqrt{8/9}L^u$, for $\alpha = \pm 1$. More generally, Appendix A.1 shows that
any membrane conductance heterogeneity decreases $L$ in comparison with the corresponding uniform case, making the cable electrotonically more compact. However, this does not necessarily imply that voltage attenuation is reduced in these more compact cables (see Discussion).

**Synapses as input**

Realistic synaptic inputs involve conductance changes, which implies that the synaptic current nonlinearly depends on the synaptic conductance change $G_{syn}$. The difference in input resistance between the slope and the uniform case (Fig. 2B) leads to a different degree of local voltage saturation in these two cases, and consequently, to differences in the amount of current generated by the synapse.

The voltage response at $x = 0$, attributable to the steady-state activation of a synapse at location $x$:

$$V(0) = g_{syn} E_{syn} \left( \frac{G_{m}(x)}{G_{m}(x) + G_{in}(x)} \right) R_{s,0}$$

(6)

where $E_{syn}$ is the reversal potential of the synaptic current, and $G_{in}(x) = 1/R_{m}(x)$ is the input conductance at location $x$ before the activation of the synapse, as computed in Figure 2B. Note that the right side of the equation is simply composed of the local input current produced by the synapse, multiplied by the transfer resistance from the input site to the soma. The derivation of this equation is given in Appendix C.2.

Figure 4A shows the voltage at $x = 0$ in response to the steady-state activation of an excitatory synapse in the maximal slope case and the uniform case (bottom two curves). As a reference, the voltage response to a DC current input, $I = g_{syn} E_{syn}$, is also shown (top two curves). This is the current that the synapse would generate if the synaptic current were not limited by saturation, and this is, therefore, the upper bound on the synaptic current that is actually generated. The corresponding curves are then proportional to $R_{s,0}$ (compare with Fig. 2A). As expected from synaptic saturation, $V(0)$ in both the uniform case and slope case is smaller for the conductance input case compared with the corresponding current input case. Still, as for the current input, the soma voltage is larger in the slope case for all input locations (Fig. 4A, two bottom curves). To highlight the effect of the nonuniform membrane conductance in the case of synaptic inputs, we have plotted in Figure 4B the benefit (the relative change in the soma voltage) of having a maximal slope membrane conductance rather than a uniform membrane conductance. The reference case of a DC current input, $I = g_{syn} E_{syn}$, is also shown (current input curve); the corresponding curve is independent of $g_{syn}$ (because of linearity) and is given by $(R_{s,0} - R_{m,0})/R_{m,0}$.

All of the curves in Figure 4B intersect at $X^u = 0.57$. At this site, the input resistance in the slope case is the same as in the...
uniform case (Fig. 2B), and the difference between the two cases (which is also independent of $g_{syn}$) is attributable only to differences in the transfer resistance, $R_k \cdot \alpha$ (Eq. 6). Most notable is that the relative change in the soma voltage is always positive (and is independent of $E_{syn}$). This means that when the input is a synaptic conductance change, the nonuniformity in the membrane conductance results in an increase of the soma voltage compared with the uniform case, as was also the case for current input. For input locations more distal than the point of intersection, the current input case sets a lower bound for this effect, and the benefit of having a sloped membrane conductance grows with increasing $g_{syn}$, eventually reaching a limit (Fig. 4B, dashed curve). In contrast, for proximal input locations, the current input case sets an upper bound for the effect of the sloped membrane conductance. The benefit of having a slope conductance is smaller when the input is a synapse rather than a current input, and this benefit decreases with increasing $g_{syn}$.

These results can be explained as follows. First, in the slope case, more synaptic current is generated at leakier distal inputs sites because the local voltage saturation is reduced as a result of the lower input resistance at these sites (Fig. 2B). The opposite is true for proximal input sites. Second, the transfer resistance is always larger in the slope case compared with the corresponding uniform case (Fig. 2A). All in all, distal synaptic inputs benefit twice from having sloped membrane conductance. For proximal synaptic inputs, however, there is a competition between the reduction of the synaptic current on the one hand (caused by stronger saturation) and the increase in the transfer resistance on the other hand. Because the latter dominates, synapses are always more effective in producing a larger soma voltage when $G_{m}$ linearly increases with distance. Distal synapses can exploit this membrane nonuniformity more effectively, because of a decreased synaptic saturation, than do proximal synapses. As a consequence, the largest relative benefit is obtained for weak proximal synapses and for strong distal synapses.

### Dendritic trees

The previous sections dealt with cylinders of constant diameter with both ends “sealed.” There are two main difficulties in extending the insights gained in these sections to dendritic trees with realistic geometries. The first arises from the “sealed ends” assumption and the second from the assumption of constant diameter. Real dendrites are complex branching structures with frequent changes in diameters, and although sealed end boundary condition is usually accepted for the termination of distal dendritic arbors, this condition is generally inappropriate at the proximal end of dendrites, toward which the synaptic current flows. At this end, the impedance load attributable to the soma and the other dendritic trees emerging from it may result in “leaky” boundary conditions. In the following, we will separately deal with each of these issues and then explore their interplay by considering relevant models of reconstructed neurons.

#### Boundary conditions

When the synaptic current flows from the “input dendrite” toward the soma, the steady-state boundary condition at the soma is given by the leak conductance, $G_{leak}$, which is the sum of the individual input conductances of all the other dendrites, each taken alone, plus the input conductance of the somatic membrane (Rall, 1959). The “leakiness” of the boundary condition can then be quantified by the parameter $B = G_{leak}/G_{syn}$, where $G_{syn}$ is the input conductance of a semi-infinite cylinder with the same diameter and specific properties as the “input” dendrite. $B = 0$ is the sealed end boundary condition (no leak through the termination), and $B = \infty$ is the “killed end” condition. Large $B$ values indicate a large “leak” at the boundary.

In Table 2, $B$ was computed for three reconstructed dendritic trees shown in Figure 7, assuming uniform membrane resistance of 20,000 $\Omega \cdot \text{cm}^2$ (fourth column) and 50,000 $\Omega \cdot \text{cm}^2$ (fifth column). The values obtained span two orders of magnitude, from 0.1 (in Purkinje cells) to more than 10 for the (thin) basal dendrites of hippocampal and neocortical pyramidal cells. For these basal dendrites, the soma and all other dendrites impose a large conductance load. The boundary conditions for the apical tree of these two cells are less leaky.

How do leaky boundary conditions affect the results obtained thus far? Figure 5A depicts the effect of varying boundary conditions at $x = 0$ on the transfer resistance $R_{k0}$. When the proximal termination becomes leaky ($B = 1$), the transfer resistance curves in the uniform case and in the slope case intersect; the intersection point moves closer to the proximal end as $B$ increases. In contrast to the sealed end case, in these cases distal current input gives rise to a smaller somatic voltage in the slope case compared with the uniform case.

The attenuation factor, however, remains smaller in the slope case (Fig. 5B), even with large $B$ value, although the difference between the two cases progressively diminishes as the leakiness at the termination increases. The results of Figure 5A, B demonstrate that, with increasing $B$ values, the longitudinal current flow in the cylinder becomes more and more dominated by the leakiness at the boundary. Consequently, the effects of the membrane nonuniformity become progressively less significant. This effect is further demonstrated in Figure 5C. Here a DC current is injected to the distal end of the cylinder, and the axial current $I_{ax}$ (given by

| Cell type              | Dendrite | $B_{20,000}$ | $B_{50,000}$ | d (µm) | max $I_{syn}$ |
|------------------------|----------|--------------|--------------|--------|---------------|
| Layer V pyramidal      | Apical   | 0.96         | 0.47         | 8.4    | 2.6           |
|                        | Basal ($n = 10$) | 12.1 ± 8.7   | 6.7 ± 4.8    | 2.5 ± 1 | 0.66          |
| Hippocampal CA1        | Apical   | 1.6          | 1.1          | 2.75   | 3.65          |
|                        | Basal ($n = 6$) | 8.3 ± 4.4   | 6.2 ± 3.2    | 1.3 ± 0.6 | 1.7          |
| Cerebellar Purkinje    | Apical   | 0.11         | 0.07         | 7.2    | 0.36          |

Table 2. Boundary condition ($B$) at the soma for the dendritic trees in Figure 7.

$B$ values were computed at the point of connection between the soma and the dendrite indicated (e.g., the apical dendrite). Computations were performed assuming uniform $R_{m,\alpha}$ of 20,000 $\Omega \cdot \text{cm}^2$ (third column) or 50,000 $\Omega \cdot \text{cm}^2$ (fourth column). In both cases $R_k = 200 \Omega$. For each dendrite $B = (2/R_k) \cdot \sqrt{\pi R_k G_{m,\alpha}}$, where $d_2$ is the diameter of the dendrite at the connection point with the soma (fifth column), and $R_{m}$ is the input resistance of the neuron at the soma when the dendrite is removed. The last column denotes the longest electrotonic path, from the soma to a dendritic terminal, for the uniform case ($R_{m,\alpha} = 20,000 \Omega \cdot \text{cm}^2$).
in the corresponding uniform case. This is true for this specific distal input location and is the result of the larger current loss (shunt) through the leakier distal membrane in the slope case. One may erroneously conclude from this figure that the charge transfer from \( x \) to the soma \((x = 0)\) is always smaller in the slope case for all input locations and boundary conditions. However, for many input locations, \( V_{\text{soma}} \) is larger in the slope case (Fig. 5A), which implies that the charge transfer to the soma is then larger than in the uniform case. This is true because \( I_{\text{soma}} = V_{\text{soma}}G_{\text{in}} \), and \( G_{\text{in}} \) is the same in the uniform case and the slope case (same \( B \) value). Hence, the charge that reaches the soma can be easily derived by scaling Figure 5A by the appropriate \( G_{\text{in}} \) value that corresponds to a given value of \( B \). Then, for a given \( B \) value and input location \( x \), the axial current that reaches the soma is larger in the slope case if \( R_{\text{s,0}} > R_{\text{c,0}} \).

In the figure above we imposed the same leak conductance \((G_{\text{in}})\) at the boundary for both the slope case and the uniform case. However, if we impose the slope condition on a realistic dendritic tree, we expect that for each dendrite, \( B \) will be different from its value in the corresponding uniform case. This is because the input resistance at the somatic end is increased in the slope case compared with the corresponding uniform case (Fig. 2), and thus \( G_{\text{in}} \) into this dendrite is decreased. This decreases \( G_{\text{in}} \) for all the other dendrites (and thus the \( B \) value becomes smaller). Note that Figure 5A also shows that in both the uniform and slope cases, the smaller \( B \) is, the better the voltage transfer from the dendrites to the soma. This means that a dendrite with a slope conductance gradient will impose a lesser conductance load to the other dendrites stemming from the soma, compared with a dendrite with uniform membrane conductance. Signal transfer from these other dendrites to the soma will then be improved because the boundary condition at their somatic end will be less leaky. To assess what are the effects of this interplay between conductance gradients and boundary conditions, it is necessary to consider reconstructed neurons (see below).

**Branching**

The results derived above for cylinders cannot be readily extended to branching structures. Consider the simple case of a branched cable consisting of a father branch and two daughter branches, one thick and one thin, such that this structure is equivalent to a single cylinder in the uniform case (Rall, 1959). This implies that the thick daughter branch is physically longer than the thin daughter branch. Now consider the maximal slope case in which the specific membrane conductance increases with a constant slope per unit length from the proximal end of the father branch to the distal terminals of the daughter branches. This case is analytically handled in Appendix A.2 and illustrated in Figure 6. Because the thick branch is physically shorter (and the conductance is growing as a function of distance from the soma), the total membrane conductance in this thin and shorter branch is smaller than in the thicker sibling branch, and the membrane at its termination is less leaky. Indeed, most of the membrane conductance is now allocated to the thick branch.

One consequence of this asymmetry in the allocation of ion channels is that the branched structure is no longer equivalent to a single cylinder in the slope case; the thin branch is now electrically shorter than the thick branch. Another consequence is that the total conductance along a given path from the soma to some terminal is not the same in the uniform case and the nonuniform case. Thus, when comparing the electrotonic properties of this structure between the uniform and slope case, the
A branched dendritic structure is generally not equivalent to a cylinder even in the uniform case, and in principle, the transfer resistance from distal sites to the soma may then be smaller in the slope case than in the uniform case. An example is shown in Figure 6B in which the thin branch is physically longer than the sibling thick branch (see inset), so that the specific conductance of its terminal is larger than in the other branch. The transfer resistance for distal input locations on this branch steeply decreases with x, and it reaches lower values in the distal part (arrow) in the slope case compared with the uniform case. This effect is observed in the models of reconstructed dendritic trees analyzed below, where some of the distal branches are much longer than others.

It is important to note that other rules for increasing membrane conductance could be implemented on dendrites displaying geometrical nonuniformities such as branching, tapering, or flare. For instance, a reasonable alternative is that the specific membrane conductance increases with distance as a function of the membrane area rather than with distance (i.e., \( G_m(x) \sim \pi \int_0^l d(z) dz \)). Another possibility is that the membrane conductance increases as a function of the electrotonic distance from the soma (i.e., \( G_m(x) \sim \int_0^x d(z) \lambda^0(z) \)). In the latter case, a branching structure that is equivalent to a single cylinder in the uniform case is also equivalent to a cylinder in the nonuniform case.

Reconstructed trees

The net effect of membrane nonuniformity on the transfer resistance for three reconstructed dendritic trees (all known to have active, and spatially nonuniform, dendritic properties) is shown in Figure 7. In all cases, \( G_m = 20,000 \text{ S/cm}^2 \). In the slope cases, the membrane conductance linearly increases with the physical distance from the soma, but the total amount of conductance was kept equal to the uniform case (such that the average value of membrane conductance, \( \overline{G}_m \), computed with respect to membrane area, was equal to \( G_m \)).

We first consider the case of the cerebellar Purkinje cell (Fig. 7A) where only one profuse dendritic tree stems from the soma. As can be seen, the transfer resistance from most of the dendritic locations to the soma is larger in the slope case (heavy line). For this neuron, the boundary condition at the soma end is close to a sealed end (Table 2); this is a favorable condition for the enhancement of signal transfer by the conductance gradient (see above). However, notice that because this cell is electrically very compact, the net effect of membrane nonuniformity is rather small (Fig. 2A, \( L^u = 0.5 \); Table 2, last column).

We next consider a neocortical pyramidal cell (Fig. 7B) where a clear distinction exists between short basal dendrites and an apical tree, with a large and thick trunk and a distal tuft. In the slope case (heavy line), \( G_m(x) \) linearly increases with the same slope in all dendrites. In the basal trees, the transfer resistance is almost twice as large in the slope case compared with the uniform case. \( R_{x,0} \) is also larger in the slope case for the main trunk of the apical tree. In the apical tuft, however, \( R_{x,0} \) is lower compared with the uniform case. This is because of the combination of the higher total \( G_m \) of the apical dendrite in the slope case (producing a larger shunt, on the average) and the conductance gradient along this dendrite that produces a strong local shunt at distal sites. As a consequence, distal inputs are less efficient in this case compared with the uniform case. Note that this example encapsulates many of the points discussed so far. The boundary condi-

Figure 6. Signal transfer in nonuniform branching geometries. Transfer resistance \( R_{x,0} \) is plotted as a function of input location, x, in a branched structure. The uniform case (dotted lines) and the maximal slope case (continuous line), in which \( G_m(x) \) linearly increases with the same slope along all the branches, are shown. The geometry of the branched structure (inset) is as follows. A, \( \ell = 500 \mu m \) and \( d = 4 \mu m \) for the father branch; for the thin daughter branch, \( \ell = 316 \mu m \), \( d = 1.6 \mu m \), and for the thick daughter branch, \( \ell = 453 \mu m \), \( d = 3.3 \mu m \). In the uniform case this structure is equivalent to a single cylinder of length \( L^u = 1 \). B, Same as in A, but the thin daughter branch is 200 \( \mu m \) longer (see inset). In both A and B the constraint of a fixed total membrane conductance (Eq. 2) was imposed. This yields a slope conductance of 1.042E−5(S/cm^2)/100 \( \mu m \) in case A and 0.967E−5(S/cm^2)/100 \( \mu m \) in case B.

Insights gained from a single cylinder are not directly valid because the constraint of the total amount of conductance imposed on the whole structure is not satisfied for each path separately.

Because of the asymmetry in the electrotonic properties of the two daughter branches in the slope case, the transfer resistances from the terminals of these branches to the soma are no longer equal as they were in the uniform case. Figure 6A shows that \( R_{x,0} \) from distal sites in the thin branch is larger than \( R_{x,0} \) from the corresponding sites of the thick branch. Still, \( R_{x,0} \) is larger in the slope case than in the uniform case, whatever the input location x is, as was the case for single cylinders. This is true even when the asymmetry between the two daughter branches is very strong. In that case, the diameters of the father branch and of the thick daughter branch are nearly equal, and these two branches almost constitute a single cylinder of constant diameter, whereas the second (very thin and short) daughter branch now has negligible membrane conductance.

The geometry of the branched structure is equivalent to a single cylinder in the nonuniform case. The uniform case (dotted lines) and the maximal slope case are nearly equal, and these two branches almost constitute a single cylinder of constant diameter, whereas the
tions at the soma end are leaky (Table 2), there is a reallocation of conductance from the shorter basal dendrites to the longer apical tree, and there are long paths in this tree where the specific membrane conductance becomes very high.

The same general behavior also holds true for the hippocampal CA1 neuron (Fig. 7C), but the advantage of having slope conductance is also apparent at distal apical arbors. We should emphasize that the results shown in Figure 7 are for steady current input; with a synaptic conductance change, this effect is expected to be even larger, especially in thin arbors in which local synaptic saturation may become significant (Fig. 4B).

The enhancement of signal transfer in basal dendrites of pyramidal cells on Figure 7B, C is caused mainly by the reallocation of the conductance in the slope case from the short basal tree to the long apical tree. This effect is analyzed on a very simple model in Appendix A.2. To control for the effect of conductance reallocation from one sub-tree to the other (e.g., basal to apical), we model in Figure 7D the case in which each of these sub-trees conserves its own total conductance as it has in the uniform case. Note that this implies that the slope per unit length is different between the two sub-trees (see inset). The enhancement of voltage transfer in the basal tree is now drastically reduced compared with Figure 7B. For the same reason, the difference between the slope and uniform cases in the hippocampal neuron (Fig. 7C) is less dramatic than in the layer 5 pyramid. Indeed, the asymmetry between the apical and basal tree of the CA1 neuron is smaller, and thus, the reallocation of \( G_m \) is less marked in this case.

As shown in Table 2, the \( B \) values at the soma are quite different between the basal and apical tree. For the basal trees, a large conductance load is imposed by the soma and apical dendrite (\( B \sim 10 \)), especially when conductance reallocation to the apical tree is allowed (Figs. 7B, C). Because the effect of gradients in membrane conductance diminishes with large \( B \) values, one expects that nonuniformity in membrane conductance will have a relatively small effect for the basal tree. This was verified by comparing the case where the conductance linearly increases in the basal tree with the case in which the same total conductance in the basal tree is uniformly distributed (results not shown). For the apical tree, however, the soma and the basal tree impose a much smaller conductance load (\( B \) values are smaller), especially if conductance reallocation between the basal and apical trees is allowed. This explains why gradient membrane conductance can improve signal transfer in the apical tree.

**Transient current injections**

In the above we dealt with the steady-state case, which is also applicable to the behavior of the time-integral of transient voltages (Barrett and Crill, 1974; Rinzel and Rall, 1974). In this section we consider transient current injection \( I(x, t) \) and restrict ourselves to the case of a single cylinder.

The solution of the time-dependent cable equation, which linearly depends on the current input \( I(x, t) \), can be obtained by convolving this current input and the Green function \( g_i(x, t) \) of the problem. \( g_i(x, t) \) is the voltage response at \( x \), \( t \) to a current pulse at time \( t = 0 \) and location \( x = x_m \). In the uniform case, \( g_i(x, t) \) can be written as an infinite sum of functions, which exponentially decay with time:

\[
g_i(x, t) = \sum_{n=0}^{\infty} c_j \delta(x - x_m) e^{-jnt},
\]

The coefficients, \( c_j \), are given by the expansion of the current impulse \( I(x, t) = \delta(x - x_m) I(t) \) on the set of orthonormal functions \( \delta_j \), so that \( c_j = \frac{\delta_j(x_m)}{\ell} \) (Rall, 1969). For sealed ends
boundary conditions, the system time constant $\tau_0$ is equal to the membrane time constant, $\tau_m^u = C_m/G_m$, and the associated function, $\psi_0 = 1$, is spatially constant. The smaller “equalizing” time constants $\tau_1 > \tau_2 > \cdots$ are given by:

$$\tau_j = \frac{\tau_m^u}{1 + \left(\frac{j \pi}{L}\right)^2}$$

The associated functions $\psi_j = \cos(j \pi c/\ell) / \sqrt{2}$ are delocalized over the whole interval $[0, \ell]$; only their phase varies with the position $x$, whereas their amplitude remains constant. This stems from the uniformity of the cable that precludes that any particular spatial location be privileged.

Figure 8 (left) displays the voltage profile along a cable of intermediate electrotonic length $(L^u = 1.5)$ at successive times, after a brief current injection. It highlights the striking differences between the maximal slope case and the uniform case. At time $t = 1$ ms, after the onset of the input current, the voltage profiles are essentially similar in the uniform case and maximal slope case (Fig. 8, $t = 1$ ms), but once the asymptotic regime is reached, where the decay of transients is governed by $\tau_m$, the voltage profile always remains nonuniform in the slope case, whereas the cable is isopotential (up to exponentially small corrections) in the uniform case (Fig. 8, $t = 20$ and $t = 80$). The behavior of the transients at right is elaborated in Discussion. This nonuniformity reflects the behavior of $\psi_0(x)$ in the slope case, which decreases from the proximal low conductance region to the distal high conductance region, as is proved in Appendix E.3. Moreover, the final relaxation phase proceeds more slowly in the nonuniform case. These differences can be understood by studying the Green function $\mathcal{G}(x)$ of the slope case.

An expansion of time-dependent solutions as a sum of orthogonal functions that decrease exponentially in time can also be derived in the nonuniform case. The time constants, $\tau_j$, and the associated eigenfunctions, $\psi_j$, are solutions of the eigenvalue problem (Eq. 5), which can be recast into the dimensionless form:

$$(L^u)^2 \frac{d^2 \psi_j(y)}{dy^2} - \left( \gamma_m(y) - \tau_m^u / \tau_j \right) \psi_j(y) = 0,$$

where $y = x/\ell$ and $\gamma_m(y) = G_m(y)/G_m$. The normalization of $\psi_j$ then becomes $\int_0^L \psi_j^*(y) \psi_j(y) dy = 1$. In the slope case, this eigenvalue problem is analytically tractable (see Appendix E). The eigenvalues and eigenfunctions depend not only on the conductance profile, $\gamma_m(y)$, but also on the electrotonic length, $L^u$, which plays the role of a diffusion constant in the above equation and controls the degree of spatial averaging. In particular, full spatial averaging occurs in the limit $L^u \to 0$, where the diffusion length $\lambda^u$ becomes large with respect to the length of the cable, so that the time constants tend to the values obtained for the uniform case, whatever the actual conductance profile along the cable is. For the benefit of the physics-oriented reader, the effects of $L^u$ are discussed in Appendix D.1 using a formal analogy with quantum mechanics.

In Figure 9A, the system time constant, $\tau_0'$, and the first two equalizing time constants, $\tau_1'$ and $\tau_2'$, are plotted in the maximal slope case and for sealed ends boundary conditions, as a function of $L^u$ in the range $0 \leq L^u \leq 10$. This range should cover all physiologically relevant situations, even when strong background synaptic activity increases the effective electrotonic length of the dendritic tree (Bernander et al., 1991; Rapp et al., 1992). In this range, the system time constant $\tau_0$ in the slope case is larger than in the corresponding uniform case. This is actually true for all values of $L^u$, and not only in the slope case, but also whenever the membrane conductance is not uniform (as proved in Appendix D.2). For the general slope case, it can be shown that conversely $\tau_0 < \tau_m^u(1 - \alpha)$ (see Appendix E.3). Moreover, $\tau_0'$ monotonically...
increases with \( L^u \). In the limit of small \( L^u \), it can be shown (see Appendix E) that:

\[
\tau_0^u = \frac{\tau_m^u}{1 - \frac{(L^u)^2}{30} + O((L^u)^4)},
\]

in the maximal slope case. In the opposite limit of large \( L^u \), \( \tau_0 \) goes to \( \tau_m^u/\min G_m(y) \), as shown in Appendix D.3 (so that it diverges to infinity in the maximal slope case, because \( \min G_m(y) = 0 \)). As a consequence of this behavior, the final relaxation of the potential, after fast equalization has occurred, proceeds more slowly in all nonuniform cases compared with the uniform case.

The equalizing time constants \( \tau_j, j > 1 \), all vanish with \( L^u \) as \( (L^u)^2 \), as in the uniform case (see Appendix E). However, at variance with \( \tau_0^u \), they are smaller in the maximal slope case than in the uniform case for small \( L^u \) values, as proved in Appendix E.2. This is in contrast with their behavior at large \( L^u \), where they are larger than in the uniform case and go to infinity with \( L^u \) (see Appendix E.2). For instance, \( \tau_1 \) is smaller than \( \tau_0^u \) as long as \( L^u < 10 \), and then becomes larger than \( \tau_0^u \), as can be seen in Figure 9A. Note also that \( \tau_j^u \) significantly departs from \( \tau_0^u \) only for large enough \( L^u \) (typically \( L^u > j^u \)). This means that, at variance with \( \tau_0 \), the time course of voltage equalization is little affected by the conductance gradient for physiologically reasonable values of \( L^u \).

In this range, differences with respect to the uniform case occur only for the first equalizing time constants and only for rather large values of \( L^u \) (\( L^u > 3 \)) and do not exceed 20%.

As in the uniform case, the eigenfunctions oscillate faster and faster as \( j \) increases, the function \( g_j \) vanishing exactly \( j \) times on the open interval \((0, \ell)\). This is illustrated in Figures 9B, C, where the first two modes, \( g_0^u(y) \) and \( g_1^u(y) \), are plotted for the maximal slope case and sealed end boundary conditions. At variance with the uniform case, they are not pure Fourier modes: they show some degree of spatial localization, their amplitude being larger in the low conductance region of the cable (Fig. 9B, C). In particular, \( g_0^u(x) \) is not a constant. For sealed end boundary conditions, it is a monotonic function of \( x \), as proved in Appendix E, reaching its maximum at the proximal end \((x = 0)\), where the membrane conductance is minimal, whereas its minimum is obtained at the distal end \((x = \ell)\) where the membrane conductance is maximal. This means that in the final stage of voltage relaxation when the voltage decays everywhere at the same rate \( \tau_0^u \), the cable is not isopotential, at variance with the uniform case. At this time, axial current constantly flows from the low membrane conductance region to the high conductance region, maintaining a voltage gradient. The nonuniformity of \( g_0^u \) increases with \( L^u \). It disappears when \( L^u \) goes to 0; the uniform solution \( g_0^u(y) = 1 \) is then recovered. For large values of \( L^u \) (\( L^u = 10 \)), \( g_0^u(y) \) develops an exponentially decreasing tail in the high conductance region (this feature is discussed in Appendix D.1). Slow voltage transients will then be observed only in the low conductance region during the final relaxation phase. We note that current injection in the high conductance region will lead to small voltage transients. The other eigenfunctions display a similar behavior as \( g_j^u(y) \), as illustrated in Figure 9C in the case of \( g_1^u(y) \). As previously noticed for \( \tau_j^u \), compared with \( \tau_j^u \), larger values of \( L^u \) are required to observe strong departures of \( g_j^u(y) \) from the cosine profiles of the uniform case. This supports the general conclusion that for values of \( L^u \) typical of dendrites (\( L^u = 1 - 3 \)), conductance gradients affect the slow asymptotic relaxation of the po-

![Figure 9. Time constants and eigenfunctions. A. The system time constant \( \tau_0 \) and the first two equalizing time constants, \( \tau_1 \) and \( \tau_2 \), are normalized to the membrane time constant \( \tau_0^u \) and plotted for a cylinder with sealed ends, as a function of the electrotonic length \( L^u \) (\( L^u \leq 10 \)). Both the uniform case (\textit{dotted lines}) and the maximal slope case (\textit{solid lines}) are shown. Explicit analytical solutions are available in the uniform case. The values in the slope case were computed by solving a transcendental equation as explained in Appendix E. B. Eigenfunction \( g_0^u \) as a function of the dimensionless variable \( y = x/\ell \) in the maximal slope case for two values of \( L^u \) (2 and 10) and in the limit \( L^u \to 0 \) (where the uniform case is recovered). \( g_0^u(y) \) was computed as explained in Appendix E. C. Same for the eigenfunction \( g_1^u(y) \). The convention that \( g_0^u(0) > 0 \) was adopted. The reader is referred to Appendix D.1 for further elaborations.](image-url)
tential but have little effect on the faster “equalization” process. Note also that steeper conductance profiles lead to a slower relaxation and a stronger localization of τ₀ (result not shown).

**DISCUSSION**

**Main results and insights**

The first general conclusion drawn from this work is that for cylindrical cables with both ends sealed, monotonic increase in $G_m$ increases the transfer resistance from any input location to the proximal end (soma) compared with the corresponding uniform case. Thus, an increasing $G_m(x)$ results in an increase of the voltage response at the soma. This effect is more pronounced when the steepness of $G_m(x)$ increases. Importantly, this improvement, caused by monotonic increase in $G_m$, is also generally valid in reconstructed passive dendritic trees, although exceptions are expected for relatively long distal dendritic arbors. When synaptic inputs (rather than current inputs) are involved, the advantage of membrane nonuniformity becomes even more pronounced (double “benefit”) for distal synaptic inputs. This is caused by the larger leakiness of the membrane at distal dendritic sites in the nonuniform case, which results in a decrease in the local synaptic saturation, and thus, with the generation of more current by these distal synapses [see also Bernander et al. (1994)].

The second general result is that any membrane nonuniformity decreases the electrotonic length ($L$) of the cable, compared with the corresponding uniform case. However, this increase in the cable “compactness” in nonuniform cylinders does not necessarily imply an increase in voltage (or charge) transfer because $L$ is not the sole parameter of importance in this case. Indeed, for both geometrically and electrically nonuniform cylinders, voltage attenuation also depends on the direction of current flow. For example, in cylinders with both ends sealed and with a monotonic increase in membrane conductance, the attenuation of voltage from the leaky end to the other end is smaller than in the reverse direction.

Another general result, which holds true for all cable structures with sealed ends (as neurons typically are), is that any membrane heterogeneity slows the final relaxation of voltage as compared with the corresponding uniform case. In other words, for any nonuniformity, the slowest time constant, $τ_u$, is always larger than $τ_u^m$, the slowest time constant in the corresponding uniform case. Thus, the final decay (the “tail”) of the excitatory synaptic potentials (e.g., the somatic EPSPs) is expected to be slower when the dendritic membrane conductance is spatially nonuniform, and this effect increases with $L$. The slowing down of the final decay attributable to membrane nonuniformity has important implications for synaptic integration and for the experimental estimation of $L$ (see below).

A general insight from studying the transient signals is that the system time constant $τ_u$ is a global quantity (i.e., it is the same everywhere, even in a heterogeneous cable). This results from the spatial averaging that takes place in the dendritic tree. Consequently, the properties of local electrical signals in dendrites are shaped, to a large extent, by the passive characteristics of the dendritic tree. Even an active local signal is strongly affected by the passive properties of the not-yet-activated adjacent regions. Thus, one cannot fully understand active phenomena in neurons without understanding the interplay between the underlying passive and active mechanisms.

Finally, we have shown that in cable structures with membrane nonuniformity and both ends sealed, there is a voltage gradient (an axial current) when the voltage relaxes according to $τ_u$, whereas in the uniform case, the whole structure is isopotential at this time. In the former case, the voltage is larger at sites with less leaky membrane conductance (namely, axial current constantly flows from these sites to the leakier regions). The experimental implication of this result is highlighted below.

**Implications for experiments: estimating $L$ and assessing membrane nonuniformity**

The theoretical results presented in this work have two direct implications from the experimental viewpoint. The first concerns the estimation of the electrotonic length, $L$, of the dendritic tree. One of the most powerful experimental outcomes of Rall’s cable theory for dendrites was his “peeling” method for estimating $L$ from the two slowest time constants ($τ_0$ and $τ_1$), which were “peeled” from experimental transients (Rall, 1969). Rall has analytically shown that, in uniform passive cylinders with both ends sealed:

$$L = \frac{\pi}{\sqrt{τ_0/τ_1} - 1}.$$  \hspace{1cm} (7)

What happens when this equation is used with the first two time constants estimated by peeling transients in the nonuniform case, namely, when $τ_0^m$ and $τ_1^m$ (in the slope case for example) are used in the above equation? We have shown above that $τ_0^m$ is larger than $τ_u^m$ and that, for relatively short cables, $τ_1^m$ is smaller than $τ_1$. Therefore, one expects that the estimated $L$ value, $L_{peel}$, in the slope case will be smaller than $L^u$ in the corresponding uniform case.

This is indeed the case as shown in Figure 10 where the $L$ value estimated from peeling transients in the uniform case is compared with the estimate obtained when voltage transients of the slope case are peeled. In the uniform case, the $L$ value estimated from peeling is very close to the actual value of $L^u$ for the full range of $L^u$ value tested (45° line, ◦). In the slope case, however, this estimate saturates for cylinder with $L^u > 2$ (○). Namely, for $L^u > 2$, $L_{peel}$ underestimates $L^u$. One could rightly argue that the estimate should be compared with the generalized electrotonic length ($L = \int_0^L dx/\lambda(x)$) rather than with $L^u$. Yet, the generalized electrotonic length is not expected to saturate (Fig. 10, ●) as does $L_{peel}$. We thus conclude that when the peeling method is used in cases with monotonic increase in $G_m$, the value obtained for $L$ may severely underestimate both the value of $L$ in the corresponding uniform case ($L^u$) as well as the value of the generalized electrotonic length of the structure. We note that because $τ_0^m$ and $τ_1^m$ are independent of input location, the estimated $L$ value is, generally, also independent of the location of current input (and voltage recording).

Another theoretical prediction that has a direct experimental relevance is the “crossing,” in the nonuniform case, of voltage transients measured simultaneously at distal and proximal locations (Fig. 8, right traces, top). As discussed above, in nonuniform cables there is a constant current flow from low-conductance regions to high-conductance regions when the voltage transient relaxes with $τ_0$. In the nonuniform case, when the input is delivered to distal leaky regions, the resultant voltage is initially larger near the input site, but at later times it becomes larger at proximal, less leaky sites. This implies a crossing of the voltage transients measured proximally and distally in the nonuniform case but not in the corresponding uniform case (where the structure is isopotential at large times), as indeed shown in Figure 8 (right traces). Crossing of transients may take place also in the uniform
case in the experimentally less likely situation where one electrode is near a sealed end boundary. At this location, charge tends to accumulate, and voltage transients relax slower than in more proximal locations. This may yield voltage crossing between faster decaying proximal transients and voltage transients measured near this boundary.

In conclusion, when voltage transients recorded at two dendritic locations cross each other, this strongly suggests that the membrane conductance is not uniformly distributed over the dendrites. Such a “cross-over” was indeed recently described in the experiments of Stuart and Spruston (1998) [see also “crossing” in Rapp et al. (1994), their Fig. 8].

**Implications on membrane nonuniformity for the input-output function of neurons**

This work analyzes the flow of current from the dendrites to the soma, showing that transfer impedance to the soma increases when relatively fewer ion channels are allocated near the soma and more are allocated distally. This gradient of ion channels also implies that the dendritic tree imposes a smaller load (current sink) on the spike mechanism in the axon compared with the corresponding uniform case. The estimate is accurate for uniform cylinders (◊), but in the slope case (○), both \( L^u \) and the generalized electrotonic lengths (●) are underestimated for cylinders with \( L^u > 2 \).

This would imply that the membrane time constant is very large, and consequently, that the resetting of the dendritic voltage is very slow. Another undesirable consequence of very low membrane conductance is that each synaptic input (conductance change) will dramatically change the effective membrane time constant and input resistance. Leaky resting membrane ensures that the effective (integration) time constant and the input resistance remain within a reasonable value even when a massive synaptic input bombards the neuron (Bernander et al., 1991; Rapp et al., 1992). Redistribution of approximately the same number of ion channels in the dendritic tree enables one to modulate signal processing in the dendritic tree, still ensuring that both the temporal characteristics of the signal and its amplitude will remain within a desired operating regime.

The present study explored the most straightforward case in which the total number of ion channels within the modeled structure remains fixed. There is, of course, the possibility that the number (and type) of effective dendritic ion channels change both very rapidly (e.g., because of the activation of synaptic-gated conductances as well as of voltage-gated ion currents with fast kinetics such as \( I_A \)) or at a slower time scale where the production of new ion channels is involved. Whatever the mechanism is, it is clear that because of its membrane ion channels, the dendritic...
tree becomes a very flexible electrical device that can dynamically modulate, at different time scales its input–output capabilities.

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