TEMPORALLY-SPECIFIC MODIFICATION OF MYELINATED
AXON EXCITABILITY IN VITRO FOLLOWING
A SINGLE ULTRASOUND PULSE

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Abstract—Single, short-duration, low-energy pulses of ultrasound were found to elicit distinct modifications of the electrical excitability of myelinated frog sciatic nerve in vitro in a window extending 40-50 ms after pulse termination. These modifications include both enhancement and suppression of relative excitability, the sequence of which generally follows one of two distinct temporal response patterns. The ultrasound pulses were focused, 2-7 MHz, of 500-μs duration, and of peak intensities of 100-800 W/cm². Total absorbed pulse energies were generally less than 100 nJ/g, corresponding to local temperature rises of the nerve trunk of no more than 0.025°C per pulse, thereby precluding bulk heating as a basis of this effect. The observed effects cannot be elicited using either a subthreshold square wave or RF electrical prestimulation, suggesting a unique form of receptivity of the nerve trunk to mechanical perturbation. We present evidence that the low-frequency radiation pressure transient accompanying the envelope of the acoustic pulse is the active parameter in this phenomenon, and postulate that it may act by the gating of stretch-sensitive channels, which have been recently reported in a variety of cell membranes. These results may demonstrate that stretch-sensitive channels in neural membrane could serve to functionally modulate neuro-electric signals normally mediated by voltage-dependent channels, a finding which could suggest new clinical applications of high peak-power, low-total-energy pulsed ultrasound.

Key Words: Axon, Excitability, Pulsed ultrasound.

INTRODUCTION

A great deal of investigation concerning the effects of ultrasound on biological systems has accompanied the emergence of ultrasound in the clinical arena as an imaging, diagnostic and therapeutic tool. Much of this work has focused on exposure dosimetry, with the aim of defining specific thresholds of biological action of ultrasound, and especially in identifying the conditions under which such fields may induce deleterious effects on a living system. Ultrasound effects on the structures of the central and peripheral nervous system have received attention accordingly, and the ability to induce irreversible histological change with sufficient intensity has been demonstrated (Fry et al. 1954; Ballantine et al. 1956; Lele 1967). At acoustic energy depositions below these levels, reversible functional effects in neural systems have been observed without apparent histological alteration, including direct stimulation of neurons in vivo (Gavrilov et al. 1973, 1977a) and in vitro (Mori et al. 1987; Gavrilov et al. 1978), suppression or blockading of action potentials (Young and Henneman 1961a, 1961b; Fry et al. 1958), and direct modification of receptor potentials in mechanoreceptors (Gavrilov et al. 1977b). Reversible effects have also been reported on the electrical characteristics of other tissue types, including electrical conduction in frog muscle (Welkowitz and Fry 1956), mammalian myocardium (Mortimer et al. 1984), and frog skin (Coble and Dunn 1976).

At the membrane level of the neuron, time constants for mechanisms associated with the generation and propagation of action potentials are typically on the order of fractions of milliseconds or milliseconds. Thus, discrimination of ultrasound effects on these mechanisms necessitates the use of appropriately brief acoustic pulses coupled with a recording system capable of examining electrical characteristics of the neural preparation at different latencies, with temporal resolution on the order of these time constants. To explore the possibility of such time-specific effects, the relative excitability of a segment of frog sciatic nerve, following its irradiation by a single, fo-
cused ultrasound pulse, was studied over a range of latencies, from 0 to 100 ms post-pulse.

METHODS AND MATERIALS

Whole sciatic nerves from pithed, 3-4 in. (snout to vent) frogs of the species Rana pipiens were excised and placed in an ultrasound exposure chamber (Fig. 1) containing Ringer's solution (mM: NaCl, 115.0; KCl, 2.1; CaCl₂, 1.8; MgCl₂, 2.0; TRIS, 1.0; glucose, 3.3). The excised nerve trunks measure 3-4 cm in length from the lower tibio-fibula to the spinal ganglia, and have a nominal diameter of 2 mm. The upper dish of the chamber has a 5-mm hole at its center over which the nerve is placed. The focused ultrasound transducer (Panametrics, model V304, f = 2 in.) in the lower portion of the chamber focuses the field to a 1-3 mm spot (half-power width) on the section of nerve exposed by the dish opening above depending on frequency and location in the z-dimension. The carrier signal of a Yaesu RF transceiver (model FT-101) triggered by a Grass stimulator (model S44) and amplified (Ameritron, model AL-1200 RF amplifier) is used to drive the ultrasound transducer. Field intensities were measured using a point hydrophone (Specialty Engineering Associates, model PVDFZ44) calibrated using the calorimetric methods of Fry and Fry (1954) and Parker (1983).

The trigger signal to the ultrasound driver is also channeled to the delayed trigger input of a second Grass Stimulator (model S44) which provides an electrical stimulus output of variable amplitude, duration, and latency. On either side of the exposed nerve segment lie two Ag/AgCl electrodes separated by approximately 5 mm. Through these electrodes, a 15-μs electrical stimulus from the second stimulator is delivered, the intensity of which is adjusted prior to each experiment to yield a compound action potential (CAP) of an amplitude of approximately half of its value at saturation. Thus a percentage of fibers contributing to the CAP can be presumed to be stimulated just over threshold, and another can be presumed to be stimulated just below threshold. Any modification of excitability by the ultrasound pulse therefore results in relatively greater or fewer numbers of fibers being stimulated over threshold, which is reflected by an increase or decrease, respectively, of the CAP amplitude relative to the no-ultrasound control. A second pair of Ag/AgCl electrodes records the CAPs approximately 1.5 cm distally, where the end of the nerve bundle exits the bath. Both the stimulus and recording electrodes were oriented away from the cut ends of the nerve trunk to minimize the contribution of any injury potentials.

Two important variations of the basic protocol were utilized in addition to that described above. In one parallel experiment, the acoustic pulse was applied to a section of the nerve bundle between the points of stimulus and recording to determine its effect on a propagating CAP initiated outside the irradiated region. The second variation of the basic protocol utilized a transducer-driven glass stylus to apply a direct mechanical prestimulus to the nerve trunk in the region which was subsequently electrically stimulated at 0-100 ms latencies. The transducer consists of a high-frequency response (3-20 kHz) loudspeaker coil driven by a 500-μs voltage pulse of square-wave form. The stylus was attached with epoxy to the transducer and positioned over the top of the nerve with a micropositioner such that light contact exists at all times. The 3-mm diameter of the stylus was chosen to involve a segment of the nerve bundle comparable to that of the ultrasound prestimulus. Sufficiently rapid mechanical response of this system was confirmed by optically tracking the actual displacement of the stylus tip with respect to the applied voltage pulse. The time course of the applied voltage can

![Fig. 1. Schematic representation of the ultrasound exposure system. Stimulus electrode 1 delivers the 15-μs electrical stimulus to the nerve segment exposed to the ultrasound pulse used in the excitability modification study. Stimulus electrode 2 delivers the stimulus to the end of the bundle to study the ultrasound effect on a propagating compound action potential. Recording electrodes measure the CAP at the end of the nerve trunk elevated above the bathing medium.](image)
thus be presumed to accurately reflect the displacement of the stylus tip during these experiments.

RESULTS

Excitability modification by ultrasound

The relative excitability of a segment of sciatic nerve, as reflected by changes in CAP amplitude, was studied over a range of latencies from 0–100 ms post-pulse. Specific temporal windows were found in which the irradiated section showed enhancement or suppression of excitability before settling back to reference level. The sequence of the excitability changes following an acoustic pulse generally followed one of two temporal response patterns. Figure 2 shows recordings of CAPs at six different latencies which illustrate excitability changes at characteristic points in samples of data from each of the two response patterns. These two distinct patterns will henceforth be referred to as early-suppression (ES)- and early-enhancement (EE)-type responses.

Figure 3 presents the envelope function of the CAP amplitudes for each of the response types in the interval of 0–100 ms post-pulse, which more completely illustrates the continuum of the excitability modification following a single acoustic prestimulus. The relatively short lifetimes of the modification (<50 ms) in both cases is notable.

Generally the response pattern of a given nerve is clearly either an ES- or EE-type response. Occasionally, nerves exhibit temporal response patterns which appear to be combinations of the two response types at nominal pulse intensities (e.g., 250 W/cm²). In such cases, we have observed that a relative reduction of the ultrasound prestimulus intensity (e.g., to 150 W/cm²) results in a new response pattern which most closely resembles the ES-type, while a relative increase (e.g., 400 W/cm²) introduces what resembles a strong superimposed EE-type response (Fig. 4). Thus the form of the excitability modification, as well as its amplitude, at times exhibits prestimulus intensity sensitivity. It is not known with certainty what determines the response type in the general case where one predominates at all prestimulus intensities.

There is some evidence that suggests a predisposition of A fibers (large-diameter, myelinated) towards the ES-type response and B fibers (small-diameter, myelinated) towards the EE response, based on experiments in which the relatively greater conduction velocity of the A fibers was used to separate its component of the CAP from that of the B fibers (Fig. 5). In such cases, A and B fiber groups respond independently to the prestimulus, with A fibers generally exhibiting ES-type response and B fibers generally exhibiting EE-type response.

Effect of frequency

The data presented thus far was acquired using pulses of 2 MHz ultrasound. The possibility of a frequency dependence of the effects was assessed by utilizing ultrasound pulses at 4 and 7 MHz, of comparable duration and range of intensities. The general form of the responses (i.e., ES- and EE-type) was found to be consistent whether elicited by 2, 4, or 7 MHz pulses. The relative efficiency of the different frequencies for inducing the effects was studied by comparing the acoustic pulse energies required to elicit a predetermined degree of enhancement or suppression at a given latency. If this comparison is made based on incident pulse energies, then high frequencies are observed to be generally more effective at eliciting a given excitability change. It is well known, however, that in this frequency region the absorption and attenuation coefficients of biological tissue increase linearly with frequency. If one therefore compares the relative effectiveness of these different frequencies based on interactive rather than incident intensities, no frequency dependence for either the form or degree of excitability modification is observed. Interactive intensity was defined as \( I_{\text{int}} = I_0(1 - e^{-2dz}) \) for the purposes of this plot, where \( I_0 \) is the incident pulse intensity (W/cm²), \( d \) is the attenuation coefficient of the nerve trunk (Np/cm MHz), \( f \) is the ultrasound frequency (MHz), and \( z \) is the nominal nerve trunk diameter. Figure 6 gives acoustic pulse intensity-duration plots for 20% suppression at the 7 ms latency of a typical ES-type response at 2, 4, and 7 MHz. The generally superimposed plots illustrate the relative frequency independence of the effects over a broad range of pulse intensities and durations. It should be noted that the location of the nerve trunk in the focal region was varied at the different frequencies such that the half-power field width (i.e., spot size) was constant in each case.

Effect of ultrasound spot size

The 3-mm spot size utilized in most of these experiments represents a defocused field condition, which was used to minimize the effect of any slight variations in the lateral position of the nerve trunk in the exposure chamber from experiment to experiment. Several additional experiments were conducted in which the acoustic spot size was reduced, while maintaining the same pulse intensity. Thus although the spatial power density is in each case the same, the total energies vary with the area of the spots, and less total energy is deposited at the smaller spot sizes.

Figure 7 shows intensity-duration plots for 20% suppression at 7 ms latency of an ES-type response at
Fig. 2. Compound action potentials (CAPs) recorded from the sciatic bundle at various latencies from 0-50 ms following application of a 500-μs, 400-W/cm², 2-MHz ultrasound pulse to the stimulus region of the axonal bundle. (a) Excitability modification of the ES-type response. (b) Excitability modification of the EE-type response. In both response plots, the first CAP of each series is elicited by a 15-μs electrical stimulus of sufficient intensity to generate a CAP amplitude of approximately half of its possible maximum in the no-ultrasound (no-u.s.) condition. The electrical stimulus was maintained at this setting for the rest of the recordings shown in the series. (Modification of excitability by the ultrasound pulse is reflected by an increase or decrease, respectively, of the CAP amplitude relative to the no-ultrasound control.) The spike preceding each CAP in this and subsequent figures is an artifact of the electrical stimulus used to generate the CAP. The temporal separation (approximately 1 ms) represents the relatively slower propagation speed of the ionic conduction underlying the CAP.
Fig. 3. Plot of compound action potential (CAP) peak amplitudes of (a) the ES-type response and (b) the EE-type response, at delays ranging from 0 to 100 ms following a 500-μs, 400-W/cm² acoustic prestimulus. Each plot is represented by peak values of approximately 50 CAPs. In each series the strength and duration of the electrical stimulus used to elicit the half-maximal CAPs are identical. Changes in CAP amplitudes at different latencies indicate the modification of excitability induced by the acoustic prestimulus pulse at t = 0 ms. The horizontal line in each series represents the reference CAP amplitude of the no-ultrasound condition. The results of a typical experiment are shown.
Fig. 4. Compound action potentials (CAPs) recorded from the sciatic bundle at the indicated latencies following application of (a) 500-µs, 150-W/cm² ultrasound pulse and (b) 500-µs, 400-W/cm² ultrasound pulse, (both 2 MHz), suggesting a prestimulus intensity sensitivity of the response pattern. Recordings are from the same nerve, with identical electrical stimuli generating the CAPs, in back to back trials. Note that the response pattern in 3(a) follows that of the basic ES-type, while at a higher prestimulus intensity (b), the excitation at $t = 2$ ms and the inhibition at $t = 14$ ms suggests the emergence of a superimposed EE-type response. A complete transition into the EE-type response was not observed in this experiment at the maximum intensity available (800 W/cm²) at the same duration.

1.25, 1.63, and 2.88 mm half-power field widths. Note that over the entire range of pulse intensities studied, the smaller spot size is more effective at eliciting a given excitability modification even though the total energy content of the pulse is less. These results clearly illustrate the importance of the spatial distribution of the acoustic pulse as an experimental parameter in this study, in addition to its intensity and duration.

**Effect on propagating CAP**

When the acoustic pulse was applied to a section of the nerve trunk between the points of stimulus and recording, a partial blockade of the propagating CAP was observed. This effect was maximized when the conditioning ultrasound pulse preceded the arrival of the CAP by 6–7 ms, corresponding closely to the peak of the first window of suppression in the ES-type response pattern. Conduction blockade at 14 ms latency was not regularly observed. No effect was observed when the arrival of the ultrasound pulse and CAP coincided, nor was any increase in CAP amplitude observed at any latency in this experiment. Intensities of the prestimuli required to elicit changes in a propagating CAP are generally four to five times greater than those required to modify excitability to the same degree at the same latency in the previous experiment.

**Direct mechanical prestimulus**

When the transducer-driven glass stylus was utilized to apply a direct mechanical prestimulus to the nerve trunk (500-µs duration) analogous excitability modifications were observed compared to those elicited using the single ultrasound pulse as the prestimulus. The temporal response patterns compared closely to those characterized earlier as ES- and EE-type responses.

The implications of this experiment are significant in assessing the relative importance of several experimental parameters, most notably the frequency dependence of the general effect, and associated thermal and cavitation mechanisms. These will be addressed in the Discussion section of this paper.
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Fig. 5. Response of different fiber groups to the ultrasound prestimulus. The two major classes of myelinated fiber can be differentiated by their relative diameters, resulting in different conduction velocities. Components of the total CAP contributed by the respective fiber groups will thus emerge as the slower component of the smaller B fibers lagging that of the larger and faster A fibers during conduction along a length of axon. In the no ultrasound recording, amplitudes of the A and B fiber component are approximately equal. The subsequent recording shows the nerve response to a 500-μs, 250-W/cm², 2-MHz acoustic pulse preceding the electrical stimulus by 7 ms. Note that the A fiber component exhibits a suppression of excitability, consistent with the ES-type response, while the B fiber component exhibits excitability enhancement, consistent with the EE-type response at this latency.

DISCUSSION

As with all phenomenological observations, direct causality between the experimental parameters and system responses cannot be assumed, necessitating careful consideration of the range of factors which may underlie the observed effects. Several modes of possible interaction of the ultrasound field with the nerve trunk are considered below.

Thermal effects

Ultrasonic fields often interact with living systems as a result of bulk heating produced by acoustic absorption. Many of the early studies reporting nerve conduction effects utilized cw- or long-pulse ultrasound, and correlated the observed changes to the accompanying temperature rise (Lehmann and Biegler 1954). In this study, although peak pulse powers are high, total energy deposition is low, less than 100 mJ/g, owing to the short duration of exposure. For a typical nerve trunk at 2 MHz, this corresponds to a general temperature rise of less than 0.025°C. Effects due to bulk heating are therefore doubtful.

The possibility of microthermal effects, due to differential heating of specific structures of the axons, was considered, as was the significance of the relatively high rate-of-change of temperature calculated during the pulse (0.025°C/500 μs), for which a theoretical basis of effect exists (Barnes, 1984). Both of these mechanisms are, however, strongly precluded by the analogous effects observed following the 500-μs direct mechanical stimulus, since acoustic absorption and, hence, heating at the frequency band of this pulse would be negligible.

Transient cavitation

At the acoustic intensities utilized in this study, small gas bubbles in the nerve tissue or medium may exhibit complex dynamical behavior, including transient cavitation characterized by violent collapse (Flynn 1964; Neppiras 1980). Very high temperatures in the vicinity of the bubbles can dissociate water vapor into the free radicals H⁺ and OH⁻, which
ACOUSTIC PULSE INTENSITY vs DURATION

TO ACHIEVE 20% ENH. AT 2, 4 AND 7 MHz

Fig. 6. Ultrasound pulse intensity-duration plots for a predetermined effect at 2, 4, and 7 MHz. Each point on the curves represents the duration required at a given intensity to achieve a 20% suppression of the half-maximal CAP amplitude at the 7 ms latency of an ES-type response. The acoustic spot size in each case was maintained at 2 mm. Note that the pulse durations are plotted as a function of interactive intensities rather than incident intensities to normalize for the different absorption and scattering coefficients for the different frequencies. The generally superimposed nature of the plots illustrate the relative frequency independence of the effect over a broad range of pulse intensities and durations.

can interact with other components, resulting in further chemical changes.

The potential role of cavitation in eliciting the observed excitability effects was thus considered. Although some transient cavitation may occur with each pulse, the similar excitability modifications which occurred following the direct mechanical stimulus, with which cavitation would not be expected, argue against a significant contribution of this mechanism.

*Stimulus coupling artifacts*

Direct effects of the ultrasound field on the stimulus electrodes and on the coupling of the stimulus current with the nerve trunk were also considered. The location of the stimulus electrodes on either side of the nerve are separated by 5 mm which places them outside the maximum field spot of 3 mm, thereby making direct effects of the ultrasound field on the electrodes unlikely. Likewise, the recording electrodes located outside of not only the field but also the bath itself are not subject to direct interaction with the ultrasound.

Coupling of the stimulus current with the nerve trunk could be affected by movement of the bathing medium, the nerve itself, or both. The sensitivity of the set-up to nerve movement was studied by varying the position of the nerve with respect to the electrodes using a micromanipulator. Movements of several millimeters in any given direction were necessary to alter the CAP amplitude to the same degree observed following the ultrasound exposure. Microscopic ex-
ACOUSTIC INTENSITY–DURATION PLOTS

FOR 20% SUPPRESSION AT 7msec OF ES-TYPE

Fig. 7. Effect of varying acoustic spot size while maintaining incident intensity. Each point on the curves represents the duration required at a given incident intensity to achieve a 20% suppression of the half-maximal CAP amplitude at the 7 ms latency of an ES-type response. The half-power field widths used were 1.25, 1.63, and 2.88 mm at a frequency of 2 MHz. Since peak intensities in each case were identical, the energy content of a pulse at a given intensity and duration decreases as the spot size is reduced. Note that over the entire range of pulse intensities studied, the smaller spot size is more effective at eliciting a given excitability modification even though the energy content of the pulse is less.

amination during and following the ultrasound pulse revealed no gross movement of the fiber bundle with respect to the stimulus electrodes.

Stimulus current coupling could also be affected by displacement of the aqueous bath which serves as the coupling medium. We have observed that movement of the bath is reflected in the electrical stimulus artifact which precedes the CAP. The relative stability of the stimulus artifact at all latencies, while the CAP amplitude changed markedly, is therefore inconsistent with what would be expected if the coupling artifact were acting. The height of the bath also was not observed to influence the experimental results.

Basis of effect

In approaching the question of identifying the basis of the observed effects, several points are notable. First is the observation of the analogous effects of the 500-μs ultrasound pulse and that of the 500-μs direct mechanical stimulus. This finding suggests that the perturbation associated with the ultrasound pulse envelope, rather than the oscillatory nature of the mechanical perturbation within the pulse, is of greatest importance. The radiation force envelope may thus be implicated as the active parameter. This hypothesis is consistent with the observed frequency independence of the effect.

Radiation force is derived from the average momentum of an acoustic beam carried per unit time past a plane normal to the propagation axis, given by

\[ \text{Momentum} = \frac{J}{c} \]
where $I_a$ is the time-average field intensity, $A$ is its cross-sectional area, and $c$ is the speed of sound in the medium. Interaction of the field with the nerve trunk causes a change in momentum, resulting in a normal force given by

$$F_{\text{norm}} = \frac{(I_{\text{abs}} + 2I_e)A \cos \theta}{c}$$

where $I_{\text{abs}}$ and $I_e$ are absorbed and scattered fractional intensities of the field incident on the nerve, $A$ is the cross-sectional area of the irradiated region, $c$ the speed of sound, and $\theta$ is the angle between the beam axis and the normal to nerve surface. This expression, although strictly true only for a continuous plane wave in a homogeneous, nonviscous fluid medium, provides a reasonable estimate for radiation force developed in this work.

The fraction of incident intensity absorbed and scattered by the nerve can be estimated by

$$I_{\text{abs}} = I_0(1 - e^{-2\alpha z})$$

$$I_e = I_0(1 - e^{-2\sigma z})$$

where $I_0$ is the time average incident intensity during the pulse, $\alpha$ and $\sigma$ are the absorption and scattering coefficients, respectively (Np/cm MHz), $z$ is the nerve thickness (cm), and $f$ the acoustic frequency (MHz).

Assuming an acoustic spot size of 0.3 cm, a nerve trunk diameter of 0.2 cm, $c = 1.5 \times 10^5$ cm/s, $I_0 = 500$ W/cm$^2$, $\alpha = 0.039$ Np/cm MHz, and $\sigma = 0.047$ Np/cm MHz, the radiation force developed during a single pulse of 2.4 and 7 MHz ultrasound is calculated to be $2.1 \times 10^{-3}$, $4.1 \times 10^{-3}$ and $7.2 \times 10^{-3}$ dynes, respectively. These forces correspond to radiation pressures across the nerve trunk of $3.5 \times 10^{-4}$, $6.8 \times 10^{-4}$, and $1.2 \times 10^{-3}$ dynes/cm$^2$.

A comparison of the mechanical forces developed by the transducer-driven stylus and those developed by the radiation pressure of the ultrasound can be made based on calculations of the total pulse energy in each case resulting in an undirectional force on the nerve trunk. In the case of the ultrasound pulse, this would correspond to the fraction of incident pulse energy which is transduced to radiation pressure, or

$$E_{\text{rad}} = (I_{\text{abs}} + 2I_e)I_{\text{pulse}}$$

Assuming the same ultrasound parameters as above at 2 MHz, the component of pulse energy translated into radiation pressure is 1.6 mJ.

In the case of the direct mechanical stimulus, the energy applied to the nerve can be estimated based on the measured electrical energy going into the transducer and the estimated conversion efficiency to mechanical energy. Typical applied electrical pulse power in these experiments was 3.1 W. Assuming a conversion efficiency of 50%, the resulting mechanical energy applied through the stylus is

$$E_{\text{stylus}} = \frac{1}{2}(P_{\text{ele}} \cdot t_{\text{pulse}}) = 0.8 \text{ mJ}.$$ 

Thus the total pulse energies directed as a mechanical force normal to the nerve trunk in both cases is seen to be comparable, further supporting the view that the ultrasound energy interacting with the nerve as radiation pressure is the active component of the pulse.

It is interesting to note that auditory nerve responses in cats to pulses of 5 MHz, 30 W/cm$^2$ ultrasound from a transducer placed against the dura matter have been reported, and attributed to radiation pressure transients accompanying these pulses (Foster and Wiederhold, 1978). The authors assumed complete absorption of the ultrasound energy in the brain tissue, and thus calculated the peak radiation pressure per pulse to be $2.0 \times 10^{-4}$ dyne/cm$^2$. Other investigators have reported human hearing sensation to 2.35 MHz pulsed ultrasound applied to the head with frequency equal to the pulse repetition frequency (Gavrilov et al. 1980). This action of ultrasound was attributed in part to modulation of radiation pressure acting on the aural labyrinth, with an intensity threshold of 7-110 W/cm$^2$ (pulse duration of 1 ms) depending on the localization of the focal zone. Unpulsed ultrasound at the same intensity failed to elicit any auditory sensation. The expectation that radiation pressure transients accompanying pulsed ultrasound may have direct action on neural tissues is thus not without precedent in the literature, although the importance of this mode of interaction of ultrasonic field and tissue has perhaps received disproportionately little attention overall.

A second important observation is that even the earliest window of effect, the excitability enhancement peaking at 5 ms in the EE-type response, is relatively slow in the context of typical neuroelectric phenomena associated with action potential generation, such as channel switching times, or even refractory periods, which generally do not extend beyond 2-3 ms in the frog sciatic fibers. A body of evidence does exist, however, for the existence of “slow” channel types or conductance states of both potassium
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(Dubois 1981; Ilyin et al. 1980) and sodium (Benoit et al. 1985) with time constants extending well into this range. Another important observation is that no emulsion of these effects could be elicited using either a subthreshold DC or a 2 MHz RF electrical prestimulus, suggesting that the mechanisms involved are uniquely receptive to mechanical stimulation, yet translate their action into electrical excitability changes.

Such translation of mechanical into electrical energy is, of course, the defining characteristic of a mechanoreceptor, in which mechanical perturbation of the nerve ending results in changes in membrane permeability to Na and K, and hence to membrane potentials. It is interesting to note in this context that these channels are thought to be different from those involved in the production of action potentials (Kuffler et al. 1984). The possibility that similar mechanisms are playing a role in the effects that we have observed with axonal segments will be considered in future studies.

Observations of membrane sensitivity to mechanical stimulus have not been limited to mechanoreceptors, however. Stretch-activated (SA) ion channels have been reported in a variety of animal cells (Guharay and Sachs 1984; Ohmori 1984; Sigurdson et al. 1987; Lansman et al. 1987; Christensen 1987), including neural membrane, where coexisting stretch-inactivated (SI) channels have also most recently been described (Morris and Sigurdson 1989). These channels are presumed to be transmembrane structures which are activated by a conformational change during membrane tension. In these investigations, single-channel response to stretch was measured by applying suction through a patch clamp electrode. Guharay and Sachs (1984) estimated a typical membrane tension T in their experiments to be 0.67 dyne/cm. Preliminary calculations which define tension T in terms of the membrane elasticity constant K4 and fractional increase in membrane area \( \Delta A/A \) resulting from the radiation pressure of the ultrasound pulse predict tensions which are comparable to those calculated by Guharay and Sachs. It should be noted that there is considerable uncertainty associated with attempting to quantify membrane tension in the bath patch clamp context (see Sigurdson et al. 1987), and the myelinated nerve case, however it does provide a starting basis for comparison. In addition, it is interesting to note that analysis of the kinetics of SA channel activity suggests a gating mechanism which involves multiple open and closed states, with time constants falling in the range of latencies where we have observed effects.

A candidate for the mechanism underlying the excitability changes observed may thus be the transient gating of SA and/or SI ion channels, whose resulting currents could serve to modulate excitability by altering threshold. Although we are unaware of any studies in which these channels have been documented at nodes of myelinated axon, we know of no evidence to preclude this, especially given the seemingly ubiquitous distribution of SA channels among the variety of cells studied.

Effect of spot size

The relatively greater effectiveness of the smaller spot sizes, even with their reduced energy content, can be perhaps explained in this context of stretch-activated neural events. The electrode geometry in this study is such that the region of the nerve trunk undergoing direct electrical stimulation is probably limited to a few millimeters, and thus any mechanism which elicits excitability changes must be active in this relatively limited region to be observed.

Radiation pressure acting on a region of membrane would be expected to result in a local displacement, which in turn would result in an increase in area or stretch of the membrane surface. SA channels are presumed to be sensitive to tension directed parallel to the membrane surface. Thus the membrane regions in which stretch-activation would be most likely to occur may be those across which the pressure gradient, and not simply the peak pressure, is maximized along its surface. The smaller ultrasound spot sizes may therefore be more effective at eliciting an observable excitability change because of the relatively larger gradient in the spatial distribution of radiation pressure in the region undergoing electrical stimulation. One implication of this finding is that a maximal effect might be observed when the degree of stretch is maximized over a distance equivalent to the spacing between the nodes of Ranvier for a particular fiber.

In the experiment in which the effect of the ultrasound pulse on a propagating CAP was studied, several points merit discussion. The observation that maximal blockade was observed when the acoustic pulse precedes the arrival of the CAP by 6-7 ms is consistent with the major suppression of excitability in the ES-type response peaking near the same latency. It is thus likely that the mechanisms underlying these two observed effects are the same.

The relatively greater ultrasound pulse intensity required to elicit comparable suppression of the propagating CAP might be explained by the inherent safety factor (typically ~5) of myelinated axon with
regard to the amplitude of an action potential arriving at a node versus that required for depolarization past threshold. The lack of a partial blockade at 14 ms latency, corresponding to the suppression of the EE-type response, may thus be the result of the inherently less sensitive nature of this particular experiment.

At this stage of the investigation it is rather difficult to approach the question regarding the precise origins of the ES- and EE-type responses. Our experiments have demonstrated both a sensitivity to the ultrasound intensity and the fiber class being observed with regard to which response type predominates. Most striking is perhaps the independent response of the A and B fiber groups when their separate components of the CAP can be resolved as distinct peaks. It is interesting to note in this context that at least two separate investigations have ranked the relative sensitivities of the different peripheral fiber groups to ultrasound as being non-uniform, with B fibers being the most sensitive, followed by C fibers, while A fibers are reported to be least sensitive (Anderson et al. 1951; Herrick 1953). Thus if the temporal response patterns (i.e., ES- or EE-type) are determined by two or more underlying membrane events, each with different thresholds and associated time constants (e.g., gating of SA potassium, SA sodium and SI potassium channels), then an ultrasound pulse of a given intensity might evoke a different temporal response in an A or B fiber due to their differing ultrasound sensitivities. This general view is also consistent with the sensitivity of the response form to the pulse intensity which was sometimes observed. We hope to resolve this question and others regarding the exact bases of the general phenomenon in future studies using single fibers.

The implications of these observations in the clinical context are difficult to completely assess until the exact modality of interaction of the ultrasound and nerve is revealed. As a general comment, however, it can be said that any means by which neural function can be predictably and reversibly modified provides a basis for use as a system of inputting and sculpting information in the nervous system. Prosthetic, analgesic, and other therapeutic uses could thus be envisioned. Specific characterization of the ionic, metabolic, and morphologic events underlying the observed excitability modifications may lead to the coalescence of these general concepts into specific clinical applications.

In summary, these studies have shown that single brief pulses of ultrasound have the capacity to elicit temporally specific modification of excitability in myelinated axons. The low total energy of the ultrasound pulses and the comparable effects seen with direct mechanical prestimuli preclude the possibility of this phenomenon having a primarily thermal basis. These transient modifications of excitability, perhaps mediated by the activation of stretch-sensitive ionic channels, suggest a potential clinical application of high peak-power acoustic pulses in the functional modulation of neuromotoric signals.

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ABSTRACT

Single pulses of focused ultrasound have been observed to significantly modify neuronal excitability in vitro for a period of 40-50 ms following pulse termination. This window of transient modification includes periods of both relative suppression and enhancement of excitability, the sequences of which generally follow distinct temporal patterns. The ultrasound pulses were focused, 2-7 MHz, nominally of 500 μs duration, and of peak intensities of 50-800 W/cm². Specific absorbed energies were less than 100 mJ/gm, which strongly precludes bulk thermal mechanisms as a basis of this effect. Our current evidence suggests that the low-frequency radiation pressure transient accompanying the envelope of the acoustic pulse is the proximal effector in this phenomenon, acting by the gating of relatively slow stretch-sensitive channels in the neuronal membrane. These observations demonstrate the potential for high peak-power, low total-energy pulses of ultrasound to functionally modulate neuroelectric signals, a finding which could suggest new prosthetic, analgesic, or therapeutic clinical applications.

Key words: pulsed ultrasound, excitability, neuron

INTRODUCTION

Extensive investigation concerning the effects of ultrasound on biological systems has accompanied its emergence in the clinical arena as an imaging, diagnostic and therapeutic tool. Much of this work has focused on exposure dosimetry, with the aim of defining specific thresholds of biological action of ultrasound, and especially in identifying the conditions under which such fields may induce deleterious effects on a living system. Ultrasound effects on the structures of the central and peripheral nervous system have received attention accordingly, and reversible functional effects in these systems have been documented, including, direct stimulation of neurons in vivo [1,2] and in vitro [3,4], suppression or blockading of action potentials [5-7], and direct modification of receptor potentials in mechanoreceptors [8]. Reversible effects have also been
reported on the electrical characteristics of other tissue types, including electrical conduction in frog muscle [9], mammalian myocardium [10] and frog skin [11].

These studies have primarily utilized long continuous exposures, or less frequently, trains of pulsed ultrasound, with limited temporal resolution, and subsequently could observe only those bioeffects which were relatively persistent. This approach is suited for mechanisms having primarily a thermal basis, but may fail to elicit or observe bioeffects which are predominantly peak-intensity related. Study of the specific potential of high peak-power, low total-energy pulses for bioeffect is particularly relevant to the currently widespread clinical use of pulse-echo imaging systems, where peak pulse intensities typically range anywhere from 10-1,000 W/cm². Intensities of these magnitudes have the potential to evoke non-linear interactions with biological tissue which would not be apparent in exposures where the same total energy is deposited less abruptly over a longer duration. Included among these are interactions of second-order such as radiation pressure and cavitation.

It has recently become evident that micromechanical properties of neural membranes may play a significant role in the excitation process, where membrane events typically have time constants of a few milliseconds or less. In particular the existence of stretch-sensitive channels has been documented in a variety of neural membranes not normally thought of as mechanoreceptors [12-17]. The potential therefore exists for interaction between ultrasonic (mechanical) perturbations of neural membrane and the events underlying overall neural excitability. To explore the possibility of such transient effects of ultrasound, the relative excitability of myelinated frog sciatic nerve and unmyelinated lobster giant axon was studied following irradiation by a single ultrasound pulse over a continuous range of latencies from 0-100 ms.

METHODS AND MATERIALS
A detailed description of the experimental design can be found in reference [18]. In brief, whole frog sciatic nerves or lobster ventral cord were excised and placed in an ultrasound exposure chamber [Figure 1]. Specially developed instrumentation allows precise control of the timing of a brief electrical stimulus applied to the nerve with respect to a burst of ultrasound preceding it. Peak intensity of the applied pulse was varied from 50-800 W/cm² (ISPPA), with a nominal duration of 500 µs, although this was varied between 100 µs and several milliseconds for some specific experiments. In the case of the frog nerve, any modification of excitability by the ultrasound pulse results in relatively greater or fewer numbers of fibers being stimulated over threshold, which is reflected by an increase
Figure 1. Schematic representation of the ultrasound exposure system. Stimulus electrode 1 delivers the 15 µs electrical stimulus to the nerve segment exposed to the ultrasound pulse used in the excitability modification study. Stimulus electrode 2 delivers the stimulus to the end of the bundle to study the ultrasound effect on a propagating compound action potential. Recording electrodes measure the frog compound action potential (CAP) or lobster action potential at the end of the nerve trunk elevated above the bathing medium.

or decrease, respectively, of the amplitude of the elicited compound action potential (CAP) relative to the no-ultrasound control. A pair of extracellular electrodes records the CAP's approximately 1.5 cm distally, where the end of the nerve bundle exits the bath. When the lobster axon is being studied, a glass intracellular electrode is used for single-cell recording, and the stimulus current varied to determine threshold.

One important variation of the basic protocol of the frog study was utilized in addition to that described above. In this experiment, a transducer-driven glass stylus was utilized in place of the ultrasound to apply a direct mechanical prestimulus to the nerve trunk in a region which was subsequently electrically stimulated at 0-100 ms latencies. The transducer consists of a high-frequency response (3-20 kHz) loudspeaker coil driven by a 500-µs voltage pulse of square-wave form. This particular experiment was conducted because early results using extended ultrasound pulses suggested the possibility that the observed excitability effect was in response to the
envelope of the applied pulse, rather than directly to the high frequency oscillations within it.

RESULTS
Excitability Modification by Ultrasound
Specific temporal windows for both frog nerve and lobster giant axon were found in which the irradiated section showed enhancement or suppression of excitability before settling back to reference level. The lobster giant axon typically responds with a suppression of excitability peaking approximately 10 ms following exposure (Figure 2). The sequence of the frog nerve excitability changes after an acoustic pulse generally followed one of two temporal response patterns, which include periods of both enhancement and suppression of excitability. Figure 3 plots the peak CAP amplitudes in the interval of 0-100 ms following exposure for both response types, illustrating the continuum of excitability modification following a single acoustic pulse. These two distinct patterns will henceforth be referred to as early-suppression (ES) - and early-enhancement (EE) -type responses. The relatively short lifetimes of the modification (τ < 50 ms) in all cases is notable.

Figure 2. Suppression of excitability of the lobster giant axon at latencies ranging from 0-50 ms following a 500 μs, 400 W/cm² ultrasound prestimulus. The horizontal line represents the relative excitability of the unexposed nerve (control). A given percentage of reduction from control reflects the relative increase of electrical stimulus needed to reach threshold at a particular latency.
Figure 3. Plot of frog sciatic nerve compound action potential (CAP) peak amplitudes of: (top) the ES-type response, and (bottom) the EE-type response, at delays ranging from 0-100 ms following a 500 μs, 400 W/cm² acoustic prestimulus. Each plot is represented by peak values of approximately 20 CAPs. In each series the strength and duration of the electrical stimulus used to elicit the half-maximal CAPs are identical. Changes in CAP amplitudes at different latencies indicate the modification of excitability induced by the acoustic prestimulus pulse at t=0 msec. The horizontal line in each series represents the reference CAP amplitude of the no-ultrasound condition. The results of a typical experiment are shown.

The general form of these excitability modifications was found to be consistent whether elicited by 2, 4 or 7 MHz ultrasound. The relative efficiency of the different frequencies for inducing the effects was studied by comparing the acoustic pulse energies required to elicit a predetermined degree of enhancement or suppression at a given latency. If one compares the relative effectiveness of these different frequencies based on interactive intensities, i.e. normalized for calculated absorbed energies, no frequency dependence for either the form or degree of excitability modification is observed.

Generally the response pattern of a given frog nerve is clearly either an ES- or EE-type response. Occasionally, nerves exhibit temporal response patterns which appear to be combinations of the two response types at nominal pulse intensities (e.g., 250 W/cm²). In such cases, we have observed that a relative reduction of the ultrasound prestimulus intensity (e.g., to 150 W/cm²) results in a new response pattern which most closely resembles the ES-type, while a relative increase (e.g., 400 W/cm²) introduces what resembles a strong superimposed EE-type response (Figure 4).
Thus the form of the excitability modification, as well as its amplitude, at times exhibits prestimulus intensity sensitivity. It is not known with certainty what determines the response type in the general case where one predominates at all prestimulus intensities.

Figure 4. Compound action potentials (CAPs) recorded from the frog sciatic bundle at the indicated latencies following application of a (upper) 500 μs, 150 W/cm² ultrasound pulse (upper) and a 500 μs, 400-W/cm² ultrasound pulse (lower), both 2 MHz, suggesting a pre-stimulus intensity sensitivity of the response pattern. Recordings are from the same nerve, with identical electrical stimuli generating the CAPs, in back to back trials. Note that the upper response pattern follows that of the basic ES-type, while at a higher pre-stimulus intensity (lower), the excitation at t = 2 ms and the inhibition at t = 14 ms suggests the emergence of a superimposed EE-type response. A complete transition into the EE-type response was not observed in this experiment at the maximum intensity available (800 W/cm²) at the same duration.

There is some evidence that suggests a predisposition of A fibers in the frog sciatic nerve trunk (large-diameter, myelinated) towards the ES-type response and of B fibers (small-diameter, myelinated) towards the EE response, based on experiments in which the relatively greater conduction velocity of the A fibers was used to separate its component of the CAP from that of the B fibers (Figure 5). In such cases, A and B fiber groups respond independently to the prestimulus, with A fibers generally exhibiting ES-type response and B fibers generally exhibiting EE-type response.
Response of different fiber groups in the frog nerve to the ultrasound prestimulus. The two major classes of myelinated fibers can be differentiated by their relative diameters, resulting in different conduction velocities. Components of the total CAP contributed by the respective fiber groups will thus emerge as the slower component of the smaller B fibers lags that of the larger and faster A fibers during conduction along a length of axon. In the no ultrasound recording, amplitudes of the A and B fiber component are approximately equal. The subsequent recording shows the nerve response to a 500 μs, 250-W/cm², 2-MHz acoustic pulse preceding the electrical stimulus by 7 ms. Note that the A fiber component exhibits a suppression of excitability, consistent with the ES-type response, while the B fiber component exhibits excitability enhancement, consistent with the EE-type response at this latency.

Direct Mechanical Prestimulus
When the transducer-driven glass stylus was utilized to apply a direct mechanical prestimulus to the nerve trunk (500 μs duration) analogous excitability modifications were observed compared to those elicited using the single ultrasound pulse as the prestimulus. The temporal response patterns compared closely to those characterized earlier as ES- and EE-type responses.

The implications of this experiment are significant in assessing the relative importance of several experimental parameters, most notably the frequency dependence of the general effect, and associated thermal and cavitation mechanisms. These will be addressed in the following Discussion section of this paper.

DISCUSSION
As with all phenomenological observations, direct causality between the experimental parameters and system responses cannot be assumed, necessitating careful consideration of the range of factors which may underlie the observed
effects. Several modes of possible interaction of the ultrasound field with the nerve trunk are considered below.

**Thermal Effects**
Ultrasonic fields often interact with living systems as a result of bulk heating produced by absorption of the acoustic energy. Many of the early studies reporting nerve conduction effects utilized cw- or long-pulse ultrasound, and correlated the observed changes to the accompanying temperature rise [16]. In our study, although peak pulse powers are high, total-energy deposition is low, less than $100 \text{mJ/gm}$, owing to the short duration of exposure. For a typical nerve trunk at 2 MHz, this corresponds to a general temperature rise of less than $0.025^\circ \text{C}$. Effects due to bulk heating are therefore doubtful.

**Transient Cavitation**
At the acoustic intensities utilized in this study, small gas bubbles in the nerve tissue or medium may exhibit complex dynamical behavior, including transient cavitation characterized by violent collapse [20,21]. Very high temperatures in the vicinity of the bubbles can dissociate water vapor into the free radicals H+ and OH-, which can interact with other components, resulting in further chemical changes.

The potential role of cavitation in eliciting the observed excitability effects was thus considered. Although some transient cavitation may occur with each pulse, the similar excitability modifications which occurred following the direct mechanical stimulus, with which cavitation would not be expected, argue against a significant contribution of this mechanism.

**Basis of Effect**
In approaching the question of identifying the basis of the observed effects, several points are notable. First is the observation of the analogous effects on the frog nerve of the 500 $\mu$s ultrasound pulse and that of the 500 $\mu$s, direct mechanical stimulus. This finding suggests that the perturbation associated with the ultrasound pulse envelope, rather than the oscillatory nature of the mechanical perturbation within the pulse, is of greatest importance. The radiation force envelope may thus be implicated as a possible active parameter. This hypothesis is consistent with the observed frequency independence of the effect.

Radiation force is derived from the average momentum of an acoustic beam carried per unit time past a plane normal to the propagation axis, given by

$$\text{Momentum} = \frac{I_a A}{c},$$

where $I_a$ is the time-average field intensity, $A$ is its cross-sectional area, and $c$ is the speed of sound in the medium. Interaction of the field with the nerve trunk
causes a change in momentum, resulting in a normal force given by

\[ F_{\text{norm}} = \frac{(I_{\text{abs}} + 2I_{\text{sc}})A\cos\theta}{c} \]  

where \( I_{\text{abs}} \) and \( I_{\text{sc}} \) are absorbed and scattered fractional intensities of the field incident on the nerve, \( A \) is the cross-sectional area of the irradiated region, \( c \) the speed of sound, and \( \theta \) is the angle between the beam axis and the normal to nerve surface. The fraction of incident intensity absorbed and scattered by the nerve can be estimated by

\[ I_{\text{abs}} = I_0(1 - e^{-2\alpha z}) \]  

\[ I_{\text{sc}} = I_0(1 - e^{-2\sigma z}) \]  

where \( I_0 \) is the time average incident intensity during the pulse, \( \alpha \) and \( \sigma \) are the absorption and scattering coefficients, respectively (Np/cm MHz), \( z \) is the nerve thickness (cm) and \( f \) the acoustic frequency (MHz).

Assuming an acoustic spot size of 0.3 cm, a nerve trunk diameter of 0.2 cm, \( c = 1.5 \times 10^5 \) cm/s, \( I_0 = 500 \) W/cm\(^2\), \( \alpha = 0.039 \) Np/cm MHz, and \( \sigma = 0.047 \) Np/cm MHz, the radiation force developed during a single pulse of 2, 4 and 7 MHz ultrasound is calculated to be \( 2.1 \times 10^{-5} \), \( 4.1 \times 10^{-5} \) and \( 7.2 \times 10^{-5} \) dynes respectively. These forces correspond to radiation pressures across the nerve trunk of \( 3.5 \times 10^{-4} \), \( 6.8 \times 10^{-4} \) and \( 1.2 \times 10^{-3} \) dynes/cm\(^2\).

A comparison of the mechanical forces developed by the transducer-driven stylus and those developed by the radiation pressure of the ultrasound can be made based on calculations of the total pulse energy in each case resulting in an undirectional force on the nerve trunk. In the case of the ultrasound pulse, this would correspond to the fraction of incident pulse energy which is transduced to radiation pressure, or

\[ E_{\text{rad}} = (I_{\text{abs}} + 2I_{\text{sc}})A_{\text{pulse}} \]

Assuming the same ultrasound parameters as above at 2 MHz, the component of pulse energy translated into radiation pressure is 1.6 mJ.

In the case of the direct mechanical stimulus, the energy applied to the nerve can be estimated based on the measured electrical energy going into the transducer and the estimated conversion efficiency to mechanical energy. Typical applied electrical pulse power in these experiments was 3.1 W. Assuming a conversion efficiency of 50%, the resulting mechanical energy applied through the stylus is

\[ E_{\text{stylus}} = \frac{1}{2} (P_{\text{elec}} \cdot t_{\text{pulse}}) = 0.8 \text{ mJ}. \]
Thus the total pulse energies directed as a mechanical force normal to the nerve trunk in both cases is seen to be comparable, further supporting the view that the fraction of ultrasound energy interacting with the nerve as radiation pressure is the active component of the pulse.

Translation of mechanical into electrical energy is the defining characteristic of a neural mechanoreceptor, in which mechanical perturbation of the nerve ending results in changes in membrane permeability to Na and K, and hence to membrane potentials. Observations of membrane sensitivity to mechanical stimuli have not been limited to mechanoreceptors however. Stretch-activated (SA) ion channels have been reported in a variety of animal cells [12-16], including neural membrane, where coexisting stretch-inactivated (SI) channels have also most recently been described [17]. These channels are presumed to be transmembrane structures which are activated by a conformational change during membrane tension. In these investigations, single-channel response to stretch was measured by applying suction through a patch clamp electrode. Guharay and Sachs in a 1984 study estimated a typical membrane tension T in their experiments to be 0.67 dyne/cm [12]. Preliminary calculations which we have done which define tension T in terms of the membrane elasticity constant $K_A$ and fractional increase in membrane area $\Delta A/A$ resulting from the radiation pressure of the ultrasound pulse predict tensions which are comparable to those calculated by Guharay and Sachs. It should be noted that there is considerable uncertainty associated with attempting to quantify membrane tension in the both patch clamp context [13], and the myelinated nerve case, however it does provide a starting basis for comparison. In addition, it is interesting to note that analysis of the kinetics of SA channel activity suggests a gating mechanism which involves multiple open and closed states, with time constants falling in the range of latencies where we have observed effects.

A prime candidate for the mechanism underlying the excitability changes observed may thus be the transient gating of SA- and/or SI ion channels, whose resulting currents could serve to modulate excitability by altering threshold. Although we are unaware of any studies in which these channels have been documented at nodes of myelinated axon, we know of no evidence to preclude this, especially given the seemingly ubiquitous distribution of SA channels among the variety of cells studied.

**CONCLUDING REMARKS**

Caution must always be exercised in the attempt to extrapolate specific in vitro findings to the much broader in vivo context. Assessment of the implications of these observations in the clinical arena must therefore remain necessarily speculative. As a general comment, however, it can be said that any means by which neural function can be predictably and reversibly modified suggests a basis for use
as a system of inputting and sculpting information in the nervous system. Prosthetic, analgesic, and other therapeutic uses could thus be envisioned. Specific characterization of the ionic, metabolic and morphologic events underlying the observed excitability modifications could eventually lead to the coalescence of these general concepts into specific proposed clinical applications.

The relevance of this study to the use of clinical pulse-echo diagnostic systems is certainly indirect, but the widespread use of these systems and the clear bioeffects which we have observed in vitro make some comments in order. Typical pulse-echo instruments utilize pulses of 0.5-3 μs durations, at repetition rates of 1-10 kHz. Peak intensities may range from 10 -1000 W/cm². The shortest pulses which our current equipment can generate, and hence the shortest used in this study, were 100 μs, about 50 times longer than typical clinical pulse lengths. However we continued to observe excitability modification at this pulse length, thus the lower limit at which these effects can occur has not yet been revealed. We plan to conduct further studies to investigate this limit.

The general finding that the observed excitability modification can be similarly elicited using a rapid direct mechanical stimulus argues against the contribution of any resonance phenomenon in this effect. This suggests that neurons possess a broad spectrum of mechanical sensitivity at the membrane level. It is not unreasonable therefore to speculate that this functional sensitivity which we have documented might play a role in the neurological response of biological systems to other rapid mechanical stimuli as well.

In closing, perhaps the broad message which can be extracted from this investigation is that the potential exists for the elicitation of unique bioeffects by high peak-power ultrasound, through mechanisms other than cavitation, arising from nonlinear interaction with tissue. Furthermore, these studies perhaps underscore the utility of tailoring the temporal resolution of the investigative tools with which bioeffects are being sought to match the time constants governing the specific function being studied. This is perhaps particularly true with neurological systems, where we have demonstrated that bioeffects need not necessarily be persistent to be of functional significance.

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References


