Tissue Resistance Changes and the Profile of Synchronized Neuronal Activity During Ictal Events in the Low-Calcium Model of Epilepsy

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Introduction

The pattern of synchronized neuronal activity in an epileptic event changes as the attack progresses. Because synchronization of neuronal discharges can be facilitated by electrical field effects between neurons (Jefferys 1981; Snow and Dudek 1984, 1986; Traub et al. 1985), we have examined the hypothesis that, during the course of the ictal discharge, there may be changes in tissue resistance which modify the efficacy of field effects and hence influence the changing profile of the electrographic seizure.

To eliminate synaptically mediated influences, we used the in vitro low-Ca2+ model of epilepsy. In this preparation, the CA1 region of the hippocampal slice, exposed to artificial cerebrospinal fluid (ACSF) containing a low concentration (0.2 mM) of Ca2+, generates spontaneous field bursts (Haas and Jefferys 1984; Jefferys and Haas 1982; Konnerth et al. 1984; Taylor and Dudek 1982, 1984a,b), which consist of a slow negative potential lasting several seconds (the “field shift”), superimposed on which there may be population spikes comprising synchronized action potentials (Andersen et al. 1971; Haas and Jefferys 1984; Lopez-Aguado et al. 2002; Varona et al. 2000). The amplitude and frequency of the population spikes change during the course of the discharge (Bikson et al. 2003a,b; Haas and Jefferys 1984; Taylor and Dudek 1984b).

Tissue resistivity increases during epileptic seizures, at least when induced in hippocampal slices in vitro by elevated potassium (Traynelis and Dingledine 1989). Measurements of intrinsic optical signal during epileptic activity in slices suggest a time-dependent shrinkage of the extracellular space fraction (Andrew and MacVicar 1994; Buchheim et al. 2000; Hochman et al. 1995), although the relationship between intrinsic optical signal and extracellular space can be inconsistent and can depend on recording conditions (Buchheim et al. 1999). Any shrinkage of the extracellular space might be expected to modify the strength of field effects between neurons (Ballyk et al. 1991; Dudek et al. 1986). The time course and size of resistivity-changes during epileptic seizures, and their relationship with the profile of neuronal discharges is at present unknown.

Because propagation of epileptic activity across slices can take <1–2 s (Haas and Jefferys 1984; Jefferys and Haas 1982; Konnerth et al. 1984; Yaaeri et al. 1983), resistance changes due to epileptic activity would be expected to develop at different times in different parts of the slice; moreover, resistivity is known to vary in different layers of the hippocampus, both in vitro and in vivo (Holsheimer 1987; Lopez-Aguado et al. 2001). In the present study, we therefore developed a technique to measure localized resistivity changes in the pyramidal layer while simultaneously recording population spikes from the same electrode to investigate how changes in resistivity, at the level of the cell bodies, correlate with changes in population spikes.

If the profile of neuronal activity is related to localized changes in electrical resistivity, it should be possible to potentiate or oppose the effects by modifying the size of the extracellular space. We induced such changes in slices by modifying the perfusate osmolarity. Changes in osmolarity are known to affect seizure threshold in man and in experimental models of epilepsy (Andrew 1991; Andrew et al. 1989; Roper et al. 1992; Traynelis and Dingledine 1989); in the present study, we...
have investigated the effect on the pattern of neuronal activity occurring within ictal events.

METHODS

Transverse hippocampal slices (400 μm) were prepared from male Sprague-Dawley rats (180–225 g; anesthetized with ketamine and medetomidine; killed by cervical dislocation). All experiments were performed under the Animals (Scientific Procedures) Act 1986 of the United Kingdom. The slices were stored, at room temperature, submerged in a holding chamber filled with Ringer’s solution (in mM) 125 NaCl, 26 NaHCO₃, 3 KCl, 2 CaCl₂, 1.0 MgCl₂, 1.25 NaH₂PO₄, and 10 glucose, aerated with 95% O₂–5% CO₂ mixture. After >60 min, slices were transferred to an interface recording chamber (35°C) containing either normal ACSF or modified ACSF.

Spontaneous activity was induced by perfusion of slices with “low-Ca²⁺” ACSF consisting of (in mM) 125 NaCl, 26 NaHCO₃, 5 KCl, 0.2 CaCl₂, 1.0 MgCl₂, 1.25 NaH₂PO₄, and 10 glucose, bubbled with 95% O₂–5% CO₂ mixture. Slices generating activity >2 mV in amplitude were accepted in this study (n = 147). Addition of 20–40 mM mamilto to low-Ca²⁺ ACSF produced hyperosmolar perfusates; for hypo-osmolar solutions, NaCl concentration was reduced to 105 and 90 mM (–40 and –70 mMΩ). The resistivity of the perfusates was measured using a Connet 2 Conductivity Meter (Hanna Instruments, Loughborough, UK). The resistivity of (normal osmolar) low-Ca²⁺ and normal ACSF was 63.3 Ω.cm. The resistivity of –40 mM low-Ca²⁺ was 71.4 Ω.cm.

Extracellular field potentials were recorded with glass micropipettes (2–8 MΩ) filled with low-Ca²⁺ ACSF and positioned in the CA1 pyramidal cell layer. Signals were: amplified and low-pass filtered (1–10 kHz) with an Axoclamp-2B or 2A (Axon Instruments) and Neurolog NL-106 and NL-125 amplifiers (Digitimer, Welwyn, UK), digitized using a Power 1401 and Signal and Spike2 software (Cambridge Electronic Design, Cambridge, UK) and analyzed with Signal and Spike2 unless stated otherwise.

A single-micropipette (plus 1 distant return electrode) technique was developed to measure local relative extracellular resistance changes (Fig. 3) using a single standard electrophysiology amplifier (1 channel of an Axoclamp). Fifty-hertz sinusoidal currents (usually ±5 nA, occasionally ±10 nA peak-to-peak) were injected into the extracellular micropipette used to record the field potentials (using the amplifier’s circuitry); the sensitivity could be adjusted by changing the injected current amplitude. The amplifier’s bridge balance was used to adjust the amplitude of the voltage oscillations (during the inter-burst period) to a convenient baseline level. The 50-Hz sinusoidal voltage + the biological signal were then recorded by the same field electrode. The 50-Hz component of the trace was then isolated, off-line, using Fourier filter and reconstruction (Autosignal, AISN Software) to identify the changes in extracellular resistance. Similar results were obtained in the time domain by fitting a 50-Hz sinusoid to the combined raw signal to isolate the component due to the applied current, to estimate changes in resistance, and, by subtraction, the biological signal (Spike2 script remsin.exe, Cambridge Electronic Design; scripts are on the web at http://www.ced.co.uk/pru.shtml, under “script library”).

To quantify tissue resistance, we used a double-barrelled theta glass micropipette electrode (plus 1 distant return/reference Ag/AgCl electrode) technique modified from Lopez-Aguado et al. (2001); here brief (0.2 s) pulses were injected into one barrel, and the voltage change was measured at the other barrel. The ratio of voltage change measured in the slice to the voltage change measured in the bath perfusate, multiplied by the known resistivity of the bath perfusate, was used to calculate resistivity. The resistivity of the perfusates was established in separate experiments (see preceding text). The sensitivity of the electrode could change with repeated insertions into the slice (probably due to tissue adhering to the electrode tip), so the 50-Hz voltage amplitude measured in the slice was normalized to the most recent voltage amplitude measured in the bath. With both resistance measurement techniques, the micropipette was inserted >50 μm below the slice surface to increase sensitivity and avoid the high-resistance layer of (damaged) tissue (Lopez-Aguado et al. 2001).

The main advantages of the single electrode technique were: high temporal resolution, up to twice the injected sinusoid frequency; high spatial resolution set by the electrode tip diameter (Lopez-Aguado et al. 2001); the ability to record resistance changes and endogenous electrographic seizures from a single site after appropriate signal filtering; and simplicity of implementation. The single-electrode technique was not used, in the present experiments, to make absolute resistance measurements; these were made using the double-barrelled technique.

Power spectral analysis was used to identify percentage changes in the 50-Hz signal during 0.5-s epochs. “Maximum population spike amplitude” in each burst or period was defined as the mean of the five highest amplitude spikes found; “maximum synchronization” was identified as the strongest cross-correlation of activity at two sites, separated by ~0.5 mm, during any 0.5-s epoch; “minimum population spike frequency” was defined as the lowest average rhythmic frequency recorded in any 0.25-s epoch with spikes >0.7 mV. Amplitude and duration of the slow DC shifts were measured as the mean of >5 bursts. Arrays of three to four electrodes were used to estimate propagation velocities; if the initiation zone was between two electrodes, the velocity was calculated from the delay between electrodes outside this region.

Recordings of extracellular concentration of potassium ions ([K⁺]o) were performed using double-barrelled ion-sensitive microelectrodes. One channel was backfilled with low-Ca²⁺ ACSF as the reference channel. The ion-sensing channel was backfilled with 100 mM KCl solution, and its tip was silanized and loaded with a valinomycin ion exchanger (FLUKA 60398). A calibration curve, based on the Nernst equation, was fitted to data for 1, 10, and 100 mM [K⁺]o (fixed 152.25 mM [Na⁺] background) and was used to calculate [K⁺]o, which was plotted on a linear scale.

Results are expressed as means ± SE; statistical analyses were made using ANOVA or Student’s paired t-test.

RESULTS

Profile of synchronized neuronal discharges

Incubation of slices in low-Ca²⁺ ACSF (>60 min) resulted in the development of spontaneous field bursts in the CA1 region. The average inter-burst interval was 77 ± 11s. Each low-Ca²⁺ burst (Fig. 1) was characterized by a “primary” slow negative field shift; the primary burst duration was 15.9 ± 1.4 s, amplitude was 2.9 ± 0.3 mV, and speed of propagation across the slice was 0.19–2.0 mm/s; the latter was comparable to earlier measurements (Haas and Jefferys 1984). At the end of the field burst, the DC potential returned toward baseline but was frequently interrupted by single or multiple “secondary” slow negative field shifts, which lasted 1–4 s and propagated across the slice at 10–106 mm/s, considerably faster than the primary burst.

The spatiotemporal profile of the first ~1 s of low-Ca²⁺ burst has been described in detail elsewhere (Bikson et al. 2003a). Briefly, population spikes were initially small and relatively irregular with an apparently high frequency; during the first 0.5–1 s, spike size and regularity rapidly increased and frequency decreased. (Fig. 1B left, bottom trace).

During the remainder of the primary burst, the DC potential remained stable and had superimposed rhythmic, relatively high-amplitude population spikes (Fig. 1B, middle) that could
either continue throughout the burst (in 47% of slices) or be interrupted (population spikes <0.7 mV) for periods of 2–15 s (in 53% of slices); these are respectively termed “persistent-spiking” and “intermittent bursts” (Bikson et al. 2003b). Even when present continuously, population spike amplitude varied during the burst, tending to decrease toward the middle of the burst and to increase to a maximum at the end of the burst. This terminal increase was evident when rhythmic population spikes were present throughout the final 5 s of the burst (61% of slices): the maximum population spike amplitude (see METHODS) during the first second of this period (5.0 ± 0.42 mV) was less than that during the first 5 s of the whole burst (6.7 ± 0.51 mV) and also less than that during the last 1 s of the primary burst (7.6 ± 0.61 mV; \( P < 0.0001 \) in both cases; paired t-test \( n = 46 \)); maximum population spike amplitude during the last 1 s of the burst exceeded that in the first 5 s in 67% of slices (\( P < 0.01 \)).

At the end of the primary burst, secondary bursts were seen in 33 of 75 slices (Fig. 1A, ↓). These contrasted markedly with those in the primary burst in that they started with two or three high-amplitude, high-frequency population spikes, and there was an immediate transition to high-amplitude, low-frequency spiking (Fig. 1B, right). Population spikes in the secondary bursts had, on average, a higher amplitude than those at the end of the primary burst (maximum spike amplitude during secondary bursts = 9.5 ± 0.81 mV; \( P < 0.01 \)).

Large (>2 mV) population spikes could propagate across the entire CA1 pyramidal cell layer, with propagation velocities of 56–120 mm/s. The average peak correlation coefficient of population spikes in the primary burst, recorded from two sites separated by 0.5 mm, was 0.69 ± 0.05. When a part of CA1 transiently stopped firing, population spike generation on either side of that zone persisted but became desynchronized (Fig. 2).

**Tissue resistance changes during the field burst**

Absolute resistivity was measured using the two-barrelled micropipette technique (see METHODS). In control ACSF, the average resistance in the pyramidal cell layer was 1,037 ± 250 \( \Omega \cdot \text{cm} \) (\( n = 13 \)) and in the stratum radiatum was 394 ± 121 \( \Omega \cdot \text{cm} \) (\( n = 6 \)); these values are comparable to those obtained by Lopez-Aguado et al. (2001) from slices in an interface chamber. After >60 min exposure to low-Ca\(^{2+}\) ACSF, the mean minimum resistivity measured in the pyramidal layer (immediately before the field burst) was 1,231 ± 264 \( \Omega \cdot \text{cm} \) (\( n = 15 \)); during the burst, resistivity increased to a maximum of 1,507 ± 335 \( \Omega \cdot \text{cm} \) (\( n = 15 \)). In low-Ca\(^{2+}\) ACSF, measurements were also made in stratum radiatum: at this site, preburst resistivity was 241 ± 99 \( \Omega \cdot \text{cm} \) (\( n = 3 \)), and the maximum resistivity during the burst was 245 ± 106 \( \Omega \cdot \text{cm} \) (\( n = 3 \)). Although there was considerable variability in absolute resistivity measured across slices, at a single location in any given slice, the increase in resistivity during bursts in the pyramidal cell layer was consistent and significant (22 ± 3%, \( P < 0.01 \)).
We found no significant difference in resistivity, of either the pyramidal cell layer or stratum radiatum, between slices incubated in normal ACSF and the minimum (preburst) values in slices incubated in low-Ca\(^{2+}\) ACSF.

To investigate timing of resistance changes more closely, we assessed changes in extracellular resistance in the pyramidal layer using the experimental set-up shown in Fig. 3 (see METHODS). Sinusoidal currents were injected into a single field micropipette electrode, which was simultaneously used to detect both endogenous voltage changes (i.e., electrographic seizures) and the sinusoidal voltage changes, produced by the applied current, due to any alteration in extracellular resistance; the latter depends linearly on the injected current and on the resistance of the electrode, the slice tissue, bath ACSF, and return electrode (Fig. 3A). During recording of low-Ca\(^{2+}\) bursts, the only resistance that changes is that of the tissue.

No detectable increase in tissue resistance was observed before burst initiation or during the first 0.5–1 s of the burst (this was confirmed by measuring power at 50 Hz in spectral analysis of the raw trace). During the main part of the burst, however, there was a gradual (τ ~ 5 s) increase in tissue resistance (Fig. 3B, middle); power spectral analysis showed that the maximum value was reached within 0.5 s of the end of the primary burst (when no secondary bursts followed) or within 0.5 s of the onset of the final secondary burst. The same pattern of increasing resistance was seen during both persistent spiking and intermittent spiking bursts (Fig. 3, B and C) and so did not depend on continued population spike firing. Tissue resistance decayed slowly (τ ~ 10 s) back to baseline after the termination of the burst.

Changing perfusate osmolarity, effect on population spike characteristics

Osmolarity was increased by addition of mannitol, a membrane-impermeant solute and decreased by removal of NaCl, to produce osmotic challenges ranging from +40 to −70 mOsm (see METHODS). In 6 of 31 slices tested, increasing osmolarity by 30–40 mOsm abolished spontaneous bursting, similar to the results of Dudek et al. (1990). In the remaining slices, however, bursts were still present, albeit with an increase in inter-burst interval and decrease in amplitude of the negative field shift (200 ± 37 and 94 ± 2% control, respectively, during exposure to +40 mOsm). Conversely, decreasing osmolarity decreased interburst interval and increased the amplitude of the negative field shift (69 ± 10 and 140 ± 15% control, respectively, during exposure to −40 mOsm).

The main purpose of these experiments was to determine the effects of exogenous changes in osmolarity on the profile of synchronized neuronal activity within the burst; detailed results are given in Fig. 4. Reducing osmolarity increased the maximum population spike amplitude and lowered the minimum discharge frequency observed during each burst; these effects could be reversed by restoring osmolarity to normal levels.
Increased the maximum synchronization of population spike amplitudes and a higher peak potassium transient amplitude by 13% (see METHODS). Decreasing osmolarity by 40 mOsm increased pyramidal layer resistivity during the preburst and end-of-burst periods by 28 and 32%, respectively [compared with measurements made during perfusion with normal osmolar low-calcium ACSF; \( P < 0.05, n = 7 \); comparable changes have previously been described for submerged slices (Andrew et al. 1997)].

Timing of resistance changes during the burst were again monitored with sinusoidal current injection into the recording channel. During perfusion with hypo-osmolar ACSF, the transition to high-amplitude spike generation was faster (Bikson et al. 2003a), but, as in the experiments with normal osmolar ACSF, tissue resistance did not increase during the period leading up to field bursts nor during the first 500 ms of the burst.

Changing perfusate osmolarity; effect on extracellular potassium transients

It has been suggested that osmotic changes would modulate extracellular ionic transients (Andrew et al. 1989; Jefferys 1995); however, this proposition has not been previously tested directly. We measured the changes in extracellular potassium concentration associated with low-Ca\(^{2+}\) ictal bursts before and during reductions in extracellular osmolarity. We found that decreasing osmolarity (−70 mOsm) increased peak potassium transient amplitude by 13 ± 4% (\( P < 0.05 \)); its duration increased by 45 ± 18% (\( P = 0.057 \); Fig. 5; \( n = 4 \)). Reducing osmolarity by 70 and 40 mOsm (by removing NaCl) also led to the development of spreading depression in 16 of 18 and 5 of 6 slices tested, respectively, consistent with enhanced extracellular potassium accumulation (Kager et al. 2000).

**DISCUSSION**

During each low-Ca\(^{2+}\) field burst, extracellular K\(^{+}\) concentration increased, contributing to the slow negative potential recorded by field electrodes (Haas and Jefferys 1984; Yaari et al. 1983). The decreased transmembrane K\(^{+}\) gradient causes an increase in neuronal excitability. This is not manifest, however, as random firing of individual neurons; rather, neuronal discharges become synchronized, giving rise to population spikes (Haas and Jefferys 1984; Jefferys and Haas 1982; Patrylo et al. 1996; Taylor and Dudek 1984a,b). Because the Ca\(^{2+}\) concentration in the ACSF is below that needed for chemical synaptic transmission (Jones and Heinemann 1987), synchronization must be mediated by nonsynaptic mechanisms.

We have previously shown that during the early part of the field burst, population spikes develop as a result of synchronized discharges in progressively enlarging neuronal aggregates; as the number of neurons recruited into an aggregate increases, the field potential produced by that aggregate increases, resulting in still further recruitment of neurons into the discharging populations (Bikson et al. 2003a). In the present experiments, we show that, at this stage (i.e., during the 1st

**FIG. 4. Effect of changes in osmolarity on low-Ca\(^{2+}\) field bursts.** A: field bursts recorded during perfusion with standard artificial cerebrospinal fluid (ACSF; i) and perfusion with reduced [NaCl] (−70 mOsm) (ii) and after restoration of osmolarity with mannitol (iii). B: pooled results summarizing the effects of osmotic challenge on: maximum population spike amplitude (mean control = 5.9 mV); minimum population spike frequency (mean control = 26 Hz) and maximum cross-correlation of activity at two sites separated by 0.5 mm (mean control = 0.7). Slices in which activity was completely suppressed by increasing osmolarity are not included. (30 slices; bars show SE; ANOVA shows that osmolarity significantly affects each of these measures, \( P < 0.01 \).)
The steady increase in resistance during each field burst tends to increase population spike amplitude, but it is not the only factor determining spike amplitude. We have shown elsewhere that spike generation during low-Ca²⁺ field bursts can be interrupted for prolonged periods as a result of depolarization block (Bikson et al. 2003b). We suggest that the final profile of synchronized neuronal activity during the main phase of the field burst is determined by the interaction of these two opposing influences, enhanced field potentials tending to increase the number of neurons participating in each synchronized discharge, whereas depolarization block tends to reduce the number of neurons able to generate action potentials.

We found that resistance in the pyramidal layer continued to increase even during periods when neuronal firing was completely blocked. This implies that continued ion fluxes, even in the absence of action potential generation (Bikson et al. 2003b), suffice to maintain neuronal swelling with the result that enhanced field effects are still observed when and if neuronal firing returns at the end of the burst.

After the primary burst (i.e., after the DC shift starts to return toward baseline), secondary bursts were frequently seen. In marked contrast to the primary discharge, population potentials characteristically had high amplitudes from the onset (maximum population spike amplitude was 25% greater than that at the end of the primary burst) and propagated rapidly through the slice. The present experiments suggest that this results from the enhanced field effects which develop through the course of the field burst as a result of neuron swelling.

Neuron swelling and shrinkage of the extracellular space can also be induced by experimentally applied changes in osmolarity (Dudek et al.1990; Kume-Kick et al.2002; Roper et al.1992); if direct electrical interactions have an important role in shaping the neuronal discharge during a field burst, changes in osmolarity, by modifying the efficacy of these interactions, would be expected to affect the amplitude of population spikes and the correlation of spike discharges recorded across the slice. These predictions were confirmed in the present experiments (Fig. 4). In addition, we have shown (Fig. 2) that, if part of a slice transiently failed to generate population spikes, regions on either side of that area continued to generate rhythmic population potentials, but they were no longer correlated; this observation is consistent with the hypothesis that synchronization across the slice depends on the continuous conduction of field potentials through adjacent regions of CA1.

Although it is likely that the effects of osmotic change reported here are largely mediated by changes in the extracellular space (Andrew and MacVicar 1994; Andrew et al.1997; Dudek et al.1990), modifying osmolarity can have other effects on neuronal excitability (Azouz et al.1997; Huang and Somjen1997; Wan et al.1999). Higher modifications of osmolar concentration can affect membrane properties, but changes of <40 mOsm (which had a marked effect in our experiments) are reported not to affect membrane properties (Azouz et al.1997; Ballyk et al.1991). Neuronal gap junctions may contribute to synchrony (Traub et al.1985; Valiante et al.1995; Vigmond et al.1997), but it is not established that gap junctions between neurons [as opposed to those between glia (Scemes and Spray1998)] are affected by cell swelling or changes in osmolarity. Decreasing the extracellular space fraction does, however, cause an increase in extracellular potas-
sium transients that will increase excitability; in the present experiments, this mechanism would contribute to the observed changes in the amplitude of the slow field shift (Dietzel et al. 1989). The increased size of population spikes, however, requires not only that excitability of individual neurons increases but that the action potentials are synchronized. Shrinking the extracellular space enhances field effects between neurons and could account for the increase in synchronization and growth in the amplitude of population spikes that occurs, despite the opposing influence of depolarization block, during the course of ictal discharges.

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