Spreading depolarizations cycle around and enlarge focal ischaemic brain lesions

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How does infarction in victims of stroke and other types of acute brain injury expand to its definitive size in subsequent days? Spontaneous depolarizations that repeatedly spread across the cerebral cortex, sometimes at remarkably regular intervals, occur in patients with all types of injury. Here, we show experimentally with in vivo real-time imaging that similar, spontaneous depolarizations cycle repeatedly around ischaemic lesions in the cerebral cortex, and enlarge the lesion in step with each cycle. This behaviour results in regular periodicity of depolarization when monitored at a single point in the lesion periphery. We present evidence from clinical monitoring to suggest that depolarizations may cycle in the ischaemic human brain, perhaps explaining progressive growth of infarction. Despite their apparent detrimental role in infarct growth, we argue that cycling of depolarizations around lesions might also initiate upregulation of the neurobiological responses involved in repair and remodelling.

Keywords: focal brain ischaemia; stroke; spreading depression; peri-infarct depolarization; laser speckle imaging

Abbreviations: CBFIND = indicative cerebral blood flow; CSD = cortical spreading depolarization; dMCAO = permanent occlusion of a distal branch of the middle cerebral artery; ECoG = electrocorticography; LSF = laser speckle flowmetry; MCA = middle cerebral artery; PID = peri-infarct depolarization
Introduction

A group of recent studies have established that, among the factors that may affect evolution of acute brain injury in its broadest sense, there is a substantial incidence of spontaneous depolarization events that propagate across the cerebral cortex in patients who have required urgent craniotomy for traumatic brain injury (Strong et al., 2005), for aneurysmal subarachnoid haemorrhage (Dreier et al., 2006), for malignant hemispheric stroke (Dohmen et al., 2008) or for intracerebral haematoma (Fabricius et al., 2008). Such waves of cortical spreading depolarization (CSD) recently termed ‘killer waves’ (Iadecola, 2009) affect essentially all neurons and astrocytes in their path; they were originally described by Leão (1944b), who delivered a local, mild insult to the cortex to elicit the phenomenon. Leão measured CSD indirectly using extracellular recordings of (i) the associated negative direct current shift (Leão, 1947) and (ii) the ‘spreading depression’ of electrocorticographic activity (Leão, 1947). At the level of a single neuron, CSD is characterized by a depolarization to nearly zero membrane potential, which is achieved by the combination of a reduced electrochemical gradient for potassium and the opening of persistent sodium and potassium conductances (Somjen, 2004).

Work in many laboratories over the past three decades with experimental models of these four clinical conditions has repeatedly demonstrated that CSD arises spontaneously in the surroundings of freshly developing brain lesions, propagating typically at 2–3 mm/min. The term ‘peri-infarct depolarization’ (PID) (Mies et al., 1993; Hossmann, 1996) describes these events, especially for focal experimental stroke, when it occurs in tissue that is functionally disturbed and has diminished vascular and metabolic activity, but is not yet irreversibly committed to necrosis (‘penumbra’; Astrup et al., 1981). Experimental findings suggest that further growth of infarct size is proportional to the number of PIDs (Mies et al., 1993) or to their aggregate duration (Dijkmuizen et al., 1999), and that PID number is the independent, determining variable in this relationship (Busch et al., 1996). The basis for this relationship may lie in abnormal microvascular vasoconstriction in response to the depolarization (Dreier et al., 1998; Shin et al., 2006; Strong et al., 2007), and/or in one or more severe challenges to metabolism such as depletion of the tissue glucose pool (Nedergaard and Astrup, 1986; Vespa et al., 2003; Hopwood et al., 2005; Hashemi et al., 2009). The detrimental role of PIDs may be enhanced by the fact that they often emerge in clusters, each with striking periodicity (Dreier et al., 2006; Dohmen et al., 2008).

Critically, little detail is known of the spatial spread of these waves of depolarization in the periphery of lesions and thus the spatial aspect of the relation between PIDs and infarct growth. We used the perfusion response to depolarization, already described in the early work of Leão (1944a) and often investigated thereafter (Hansen et al., 1980; Lauritzen et al., 1982), as a surrogate tracker of depolarization that can now be mapped visually in real time using laser speckle flowmetry (LSF) (Dunn et al., 2001) (see Supplementary material ‘Note 1’). Direct current electrodes were also used to verify depolarization events.

Here, we describe (i) patterns of spatial spread of depolarizations for the whole circumference of an ischaemic focus in an existing stroke model—permanent occlusion of a distal branch of the middle cerebral artery (dMCAO) in rats. Unexpectedly, and of considerable interest, we observed in this model that PIDs rotated or cycled, often several times, around the small focus. In the light of these findings, we examined (ii) the spatial behaviour of PIDs after permanent occlusion of the middle cerebral artery (MCA) in cats in detail, here and in a previous dataset available to us from another laboratory (London) (Strong et al., 2007). We assessed patterns of PID spread (as seen in the restricted fraction of the ischaemic territory that can be imaged in this model) for evidence of compatibility with the concept of peri-lesion cycling. We then (iii) document PID occurrence and corresponding circular lesion growth assessed by MRI in a human patient suffering from ‘malignant’ hemispheric MCA infarction (Hacke et al., 1996) and finally (iv) compare periodicity in the occurrence of peri-lesion depolarizations in malignant stroke patients derived from another, previous data set (Dohmen et al., 2008) with equivalent experimental results in rats and cats, prompted by the suggestion that inter-depolarization interval might increase as absolute lesion volume increases in the progression from rat, to cat, to human.

Materials and methods

Adult male rats and adult female cats were used (but see also Supplementary material ‘Note 2’). The study was approved by the local Animal Care Committee and the Regierungspresident of Köln and is in compliance with the German Laws for Animal Protection.

Anaesthesia and surgical preparation in animals

Rat experiments

In six adult Wistar rats weighing 340–714 g, general anaesthesia was induced with halothane (4%) and maintained with isoflurane (0.8–1.2%) in a 70% nitrous oxide/30% oxygen gas mixture. The left femoral artery and vein were cannulated for continuous measurement of mean arterial blood pressure and serial measurement of blood gases (arterial pO2, pCO2, pH), and for i.v. administration of fluid and drugs, respectively. After tracheotomy and immobilization with pancuronium bromide (0.1 mg/kg, i.v.), artificial ventilation was started; immobilization was maintained throughout the experiment by intravenous infusion of Ringer’s solution containing gallamine triethiodide (5 mg/kg/h). Acid–base balance was controlled using sodium bicarbonate. Deep body temperature was maintained at 37°C using a heating blanket servo-controlled by a rectal temperature probe. In order to produce small cortical ischaemic lesions in the MCA territory, we used a tandem occlusion method: occlusion of a distal branch of the MCA was combined with occlusion of the ipsilateral common carotid artery (Ohta et al., 1997). In brief, the common carotid artery was first ligated. A hemicraniectomy (1.5 x 1.0 cm) was then performed over the left hemisphere, the cortex exposed and protected with warm mineral oil and imaging then started using LSF (see below). After acquiring control images, the frontoparietal branch of cortical MCA was coagulated and cerebral blood flow change was monitored for 6 h.
Cat experiments

For full details please see Supplementary material ‘Note 2’.

In a total of seven cats (Cologne) and nine cats (London), anaesthesia was induced with ketamine or medetomidine/halothane (London) and maintenance anaesthesia was established with halothane (0.6–1.2%) in a 70% nitrous oxide/30% oxygen gas mixture. The left femoral artery and vein were cannulated for continuous measurement of mean arterial blood pressure and serial measurement of blood gases (arterial pO2, pCO2, pH), and for i.v. administration of fluid and drugs, respectively. Ringer’s solution containing gallamine triethiodide (5 mg/kg/h) was infused intravenously (3 ml/h) for immobilization throughout the experiment. Acid–base balance was controlled using sodium bicarbonate. The right MCA was exposed transorbitally to be later occluded using either a device for remote occlusion or a clip (London). Deep body temperature was maintained at 37°C using a heating blanket servo-controlled by a rectal temperature probe. A mineral oil pool was established above the exposed cortex, its temperature servo-controlled to around 37°C and imaged as in experiments in rats. For maintenance of anaesthesia throughout the experiment, continuous x-clorolose infusion (5 mg/kg/h) was started following an initial bolus (60 mg/kg/15 min, i.v.).

Electrocorticography recordings in patients

First, we report on results derived from a patient who suffered from a large infarction of the MCA territory. After a decision had been made that, independent of the study, surgery was required for decompressive hemicraniectomy, we placed a subdural electrocorticography (ECoG) strip electrode containing six individual electrodes on the peri-infarct tissue of the cortex accessible from the craniectomy. Bipolar recordings were obtained from neighbouring electrodes (as described above) so that four ECoG channels were obtained; one electrode served as ground (Dohmen et al., 2008). The patient was ventilated and sedated (fentanyl plus midazolam) throughout the monitoring. Simple visual detection of spreading depolarizations and periods of ECoG depression during the online measurement was achieved by displaying the high-pass filtered ECoG signal (lower frequency limit 0.5 Hz) and the power of the high-pass filtered ECoG signal (Fabricius et al., 2006). The duration of ECoG depression until recovery (interval between depression onset and onset of restoration of activity) (Dreier et al., 2006) was measured as an indirect indicator of the tissue energy status, since restoration of ECoG activity after spreading depolarization is energy dependent.

We performed a new analysis of the raw ECoG data of the patients included in a former study on malignant hemispheric stroke (Dohmen et al., 2008), focusing now on timing of spreading depolarization events. To detect evidence for cycling (implying repeated transit of depolarization events across/along the electrode strip in one direction only), we considered for inclusion only events recurring (≥2) and spreading in one direction between the four channels (bipolar montage) on the subdural electrode strip. Original recordings from eight patients were available for this analysis. A total of 947.6 h of ECoG monitoring (mean duration of monitoring was 118.5 h ± 32.3 h) was scanned for spreading depolarizations fulfilling the above criteria. On these criteria we identified 97 intervals in seven patients. Eighty-seven of these 97 intervals (90%) showed a duration of <150 min and are displayed with this time axis in Fig. 6 for better interspecies comparison. Of 97, 10 intervals (10%) lasted >150 min.

Magnetic resonance imaging in patients

MRI was performed on a 1.5 T whole-body scanner (Philips Intera Master, Philips, Eindhoven, The Netherlands) in an axial direction (20 slices, 6 mm slice thickness, 0.6 mm interslice gap, field of view 23 cm, repetition time 3875 ms, echo time 95 ms, echo planar imaging 77).
**Statistical methods**

Values of continuous variables are given as mean ± standard deviation unless otherwise stated. In the rat dMCAO model, the difference in regional CBF\textsubscript{IND} between pre- and post-dMCAO was tested with the Wilcoxon rank sum test. In the cat proximal MCA occlusion model of stroke, inspection of the CBF\textsubscript{IND} maps as they evolved after MCA occlusion typically showed the emergence of a clear demarcation between a core area of very low perfusion, and adjacent cortex with better but not normal perfusion. Static perfusion values here were not formally analysed, since such analysis would serve only to replicate previously published work (Strong et al., 2007). Instead, we analysed the spatial behaviour of transient, spreading changes in perfusion in the peri-infarct region.

**Results**

Please view the online Supplementary Videos 1–3 in close conjunction with this text. Legends to the videos are available at the end of the Supplementary material.

**Systemic variables**

Measurements of systemic variables did not show any significant alterations between control and ischaemic conditions, determined systematically 3 h after dMCAO in rats and MCA occlusion in cats (Supplementary Table 1).

**Peri-infarct depolarizations cycle around ischaemic foci in the rat cerebral cortex following distal middle cerebral artery occlusion**

In order to image the spread of PIDs around the complete circumference of an occlusive ischaemic lesion, we performed a large craniotomy above the left cerebral cortex and occluded a distal branch of the MCA in rats (Fig. 1A): this approach proved suitable, although imaging was interrupted during performance of the occlusion. We used a tandem occlusion model with ligation of the ipsilateral common carotid artery and coagulation of the frontoparietal branch of the MCA (Brint et al., 1988). PIDs were verified with microelectrodes as intra-cortical direct current shifts. Concomitantly, LSF detected waves of regional blood flow (‘CBF\textsubscript{IND}’, indicative of absolute values of cerebral blood flow: please see ‘Materials and methods’ section) alteration coupled to the depolarizations. These CBF\textsubscript{IND} waves (surrogates for PID) allowed us to study propagation patterns of these alterations. Static analysis of circular regions of interest (diameter 2.0 mm, seven regions of interest per animal) placed on CBF\textsubscript{IND} images adjacent to and further from the coagulation point, revealed that CBF\textsubscript{IND} significantly decreased in both the two time points, namely immediately and 1 h after occlusion, compared with pre-ischaemic controls to 50% (\(P = 0.036\), Wilcoxon rank sum) in three regions of interest near to—and to 80% (non significant) in four regions of interest more distant from—this point. The effect of dMCAO is shown in Fig. 1B; here, a CBF\textsubscript{IND} threshold <15 ml/100 g/min was applied, demarcating the ischaemic zone in blue. A total of 35 CBF\textsubscript{IND} waves were observed in six individual rats. These waves originated in 29/35 cases in the border zone of the ischaemic focus. In 6/35 cases (in two individuals), waves invaded the field of view from outside it, so that an origin could not be identified. We suspect that microfocal trauma, in particular at the border of the craniotomy, may have given rise to these depolarization events. Interestingly, all waves propagated circumferentially along the border of the ischaemic focus, including those arising outside the image field. They either cycled around the focus, often multiple times (Fig. 1C and Supplementary Video 1) or they split at the point of origin and travelled then as two waves around either side of the focus until they met at the opposite side and annihilated each other (Fig. 2).

Multiple CBF\textsubscript{IND} waves, appearing often with regular periodicity in temporal clusters, were associated with a decrease in basal (inter-event) CBF\textsubscript{IND} (analysed in a circular region of interest in the border zone of the primary infarct; Fig. 1D, left) and this decrease appeared to occur stepwise with consecutive events. All animals developing such clusters of depolarization events showed comparable decreases of inter-event CBF\textsubscript{IND} (Fig. 1E, left). Similarly, the number of pixels below a CBF\textsubscript{IND} threshold value <15 ml/100 g/min increased stepwise in the same circular region of interest (Fig. 1D and E, right) documenting the growth of the ischaemic focus in response to the number of depolarization events. In the example in Fig. 1, CBF\textsubscript{IND} waves were characterized principally by hyperaemia. Detailed analysis of all 35 events in rat dMCAO revealed, however, that 17 waves were hyperaemic, 12 were biphasic with a hypoperfusion onset followed by a hyperaemic tail that was usually longer in duration and six waves consisted of monophasic hypoperfusion. In individual cases, dispersal of CBF\textsubscript{IND} waves was wide enough to allow us to position small regions of interest in both inner and outer zones of the propagation path. In such cases, examination in inner zones near the ischaemic core showed biphasic hypo-/hyperaemic or monophasic hypoperfusion CBF\textsubscript{IND} responses (that were sometimes sustained), whereas analysis of regions of interest in the outer zones always revealed monophasic hyperaemic responses (Fig. 2). Thus, depolarization-coupled CBF\textsubscript{IND} responses in rat dMCAO are compatible with a gradient of perfusion developing in border zones of ischaemic foci after arterial occlusion, as described originally, to our knowledge, by Symon et al. (1974). Depolarizations also reduce perfusion further in the border zones, as has recently been described for MCA occlusion in mice (Shin et al., 2006) and in cats (Strong et al., 2007) using LSF real-time imaging.

**Spatial patterns of spread of peri-infarct depolarizations in the gyrencephalic brain of the cat following proximal middle cerebral artery occlusion**

In the light of the above findings, we next examined patterns of spread of PIDs around larger, occlusive ischaemic foci in the gyrencephalic brain. For this, we retrospectively reviewed spatial patterns of CBF\textsubscript{IND} wave propagation in two sets of data (one set from the laboratory of A.J.S. in London and a second from this laboratory in Cologne) on MCA occlusion in the gyrencephalic...
The anatomical disposition of the ectosylvian, suprasylvian and marginal gyri, each set circumferentially and at respectively increasing distances from the Sylvian fissure and location of core ischaemia after proximal MCA occlusion, facilitates spatial and temporal analysis of spread of depolarizations (Figs 3 and 4). These figures also illustrate the restricted fraction of the ischaemic territory that we are able to image in cats undergoing MCA occlusion. Changes in perfusion in response to depolarization in these two groups of experiments have been reported previously, with the conclusion that the responses of perfusion to depolarization in the penumbra deteriorate progressively with increasing proximity to the core: monophasic hyperaemia on the medial marginal gyrus nearest collateral perfusion from the anterior cerebral artery, biphasic alteration (hypoperfusion then hyperaemia) nearer, and monophasic hypoperfusion of varying durations, sometimes very sustained, immediately adjacent to the core (Strong et al., 2007). The new analyses of spatial patterns of spread are described in detail in the Supplementary material ‘Note 2’ and the findings are summarized here. In aggregate, 142 events were analysed (60 in 9 London experiments all sampled continuously for 4 h after occlusion and 82 in 7 Cologne experiments followed for between 10 and 26.5 h after occlusion). Patterns of spread were in principle either radial (outwards from the core focus: 41% Cologne, 11% London) (Fig. 3 and Supplementary Video 2), or circumferential (spreading along the edge of the core: 59% Cologne, 89% London) along the length of one or more of the three exposed gyri, and circumferential PIDs often appeared in clusters with rather regular periodicity (Fig. 4 and

Figure 1 Repetitive cyclic CBFIND wave propagation around primary ischaemic lesion in rat cortex after dMCAO (see also Supplementary Video 1). (A) Diagram of the dorsal aspect of the rat brain. Frame (thick lines) indicates field of view for LSF. Vascular territory of distal MCA, ischaemic territory and anti-clockwise cycling (arrow) are indicated. (B) Expansion of the ischaemic lesion after repetitive propagation of eight cyclic CBFIND waves. A region of interest (white circle; 3 mm diameter) located in the boundary zone served for time course analysis of CBFIND and increase of number of pixels with CBFIND <15 ml/100 g/min (Fig. 2C and D). (C) Sequential images demonstrating multiple consecutive turns of CBFIND waves around the ischaemic lesion demarcated by dotted black line. Time after dMCAO is indicated in individual images. Please note that the surrounding of the ischaemic core turned progressively blue compared with the first image, indicating gradual expansion of the ischaemic core. (D) Time course of mean CBFIND (left) and number of pixels with CBFIND <15 ml/100 g/min (right) analysed in boundary zone region of interest (ROI) (Fig. 2B). CBFIND flow increased repetitively while basic CBFIND decreased stepwise during CBFIND wave passage. On the other side, the number of pixels with CBFIND <15 ml/100 g/min increased in the region of interest stepwise with CBFIND waves. (E) Decrease of basic CBFIND (left) and increase of the number of pixels with CBFIND <15 ml/100 g/min (right) in relation to number of CBFIND waves (numbers next to lines). Data analysed in the various rats using region of interest analysis as shown in Fig. 2D.
Supplementary Video 3). Five events in the London experiments that originated within the imaging field later in the observation period exhibited both radial and circumferential spread. In both laboratories, it seemed that events spreading radially most often occurred early in the course of post-MCA occlusion observation, whereas circumferential spread was seen later; however, this could not be verified formally and the basis for distinction between these two patterns of spread is considered in the ‘Discussion’ section. In the Cologne data, the initial core lesion was typically slightly smaller (most probably due to higher blood pressure and consequently better collateral perfusion), and this probably accounts for the higher incidence of radial spread than in London; with radial spread, a delay was invariably seen as the wave descended from suprasylvian gyrus into the marginal sulcus, before re-emerging onto the marginal gyrus (Fig. 3 and Supplementary Video 2).

The occasionally observed occurrence of circumferentially spreading events, either ‘splitting’ from a focus at the edge of the core, or entering the imaged field almost simultaneously from opposite sides as two events and propagating towards each other before colliding, is compatible with the hypothesis that PIDs in the gyrencephalic brain are capable of tracking around at least a major portion of the circumference of a focus of core ischaemia, as imaged around the entire core lesion in our rat dMCAo preparation. Whether PIDs originate at the edge of the core, or at new foci in the penumbra has been debated (Nedergaard and Hansen, 1993): the present data document origin at the edge of the core, but (temporally sporadic) circumferential events entering the field from outside may have arisen from either potential source. Given the visualization of a split of depolarization in rat dMCAO (Fig. 2), it is reasonable to suggest that dual events appearing (near)-simultaneously on the same gyrus (cat MCA occlusion) from opposite sides of the image field may reflect the same bifid pattern of origin and spread.

Peri-infarct depolarizations are associated with circular secondary growth of lesion in patients with large middle cerebral artery infarction

Clustering of depolarizations has now been also described in recent studies in patients with subarachnoid haemorrhage (Dreier et al., 2006) and ischaemic stroke (Dohmen et al., 2008) and...
prompted a comparison of human and animal data with respect to circumferential propagation. We report here on new results in a patient suffering from malignant hemispheric MCA occlusion (Fig. 5). In this patient, PIDs were recorded using an electrode strip comprising six electrodes that was positioned subdurally over peri-infarct tissue, parallel to the infarct rim. This procedure was carried out at the conclusion of hemicraniectomy performed in order to decompress massive ischaemic oedema and brain swelling. From the six electrodes of the strip, four ECoG channels (A–D) were acquired. The ECoG depression and its duration are best visualized as a reduction of the power of the high-pass filtered ECoG amplitude, which also allows simple visual detection of PID during the online measurement at the bedside. Figure 5A shows PID measurements from a 53-year-old patient, who underwent hemicraniectomy 22 h after a large MCA infarction. Monitoring was started 29 h after stroke. During the first 24 h of monitoring, PIDs were detected only in channels D and C, spreading from D to C. Both channels were acquired from the electrodes located more proximal to the infarct rim. At later time points, PIDs progressively expanded spatially along the electrode strip involving channel B and finally also channel A (Fig. 5A), the channels from electrodes positioned further from the infarct. This expansion of depolarizing events along the strip electrode over time was accompanied by a progressive decline of the EcoG, which was more pronounced in channel D, resulting finally in an almost flat signal. Additionally, we observed an increasingly prolonged duration of ECoG depression and recovery after each PID over time (Fig. 5A and B), again more pronounced in channel D than in channel A. Furthermore, repetitive PIDs occurred in this patient, often in clusters with regular periodicity (Fig. 5C), showing unidirectional propagation of multiple PIDs along the electrode strip, and therefore along the infarct rim. From the 92 depolarizations detected in this patient, 72 spread from channel D towards channel A (clockwise along the strip electrode and with regard to the edge of infarction) and 19 spread contrariwise towards channel D (counter clockwise along the strip electrode). In this same patient, we performed MRI for the first time both shortly after start (43 h, second day post-stroke) and at the end of ECoG monitoring (186 h, seventh day post-stroke), thus enabling us directly to assess alterations in infarct size occurring exactly during the period of ECoG monitoring (Fig. 5D). The follow-up MRI at Day 7, after 92 PIDs, showed substantial infarct growth, with recruitment of new cortical regions in comparison with the first MRI. Infarct growth and newly established lesions were best documented by diffusion-weighted imaging (Fig. 5D). In the surface reconstruction in particular, the concentric nature of this growth became evident. The mapping of apparent diffusion coefficients showed a reduction (hypointensity) of apparent diffusion coefficient precisely in the areas undergoing secondary ischaemia during the period of ECoG monitoring, indicating that delayed ischaemic transformation occurs in regions of secondary damage during the time of multiple PID passage.
Inter-species comparison of depolarization frequencies

In the light of the above findings we analysed and compared retrospectively the intervals between (i) unidirectionally propagating CSDs in seven patients with hemispheric stroke experiencing clusters of depolarizations (Dohmen et al., 2008); (ii) unidirectionally propagating CBF IND waves in seven MCA occlusion experiments in cats; and (iii) cycling CBF IND waves in the dMCAO model in six rats. The resulting inter-event interval histograms (Fig. 6) demonstrate that in rat dMCAO, the modal interval is considerably shorter than in cat proximal MCA occlusion, consistent with a smaller absolute lesion volume (see Supplementary material ‘Note 3’). Although intervals between consecutive depolarization events in CSD/PID clusters in human malignant stroke were rather longer than in the cat, any difference is less marked than between cat and rat, and this is addressed in the ‘Discussion’ section. This finding suggests that periodicity of occurrence at relatively regular intervals is not only a common feature of PIDs but that periodicity may result at least in many instances from circumferential multiple propagation of depolarizations around ischaemic core lesions.

Figure 4 Circumferential CBF IND wave propagation in the ischaemic boundary zone after cat MCA occlusion (see also Supplementary Video 3). (A) Diagram of the lateral aspect of the cat brain (see also Fig. 3A). Pink arrow indicates circumferential direction of CBF IND wave propagation. Temporal emergence of 14 consecutive CBF IND waves is indicated on the right. All waves (other than 12 and 13, asterisks) travelled clockwise. (B) Sequential images of waves 9 and 14 propagating circumferentially in marginal gyrus (MG) around the ischaemic focus. Time on images indicates time from onset of respective CBF IND waves. Regions of interest (on marginal gyrus; far from ischaemic core), b (on suprasylvian gyrus (SG); close to core) in first images served for time course analysis. Black bars = 5 mm. Arrow heads indicate wave front. Note that the wave front turns into hypoperfusion (blue) in 14th wave. (C) Time course of CBF IND in regions of interest a and b showing consecutive waves 9–14. Waves 12 and 13 travelled anti-clockwise. The transition from hypo- to hyperaemic appearance as the wave propagates from inner to outer boundary zone of the ischaemic focus. Note the continuous decrease of baseline CBF IND in the zone close to the ischaemic core (region of interest b). (D) Zoomed view of time course of CBF IND waves 9 and 14, showing transition from biphasic (hypo- then hyperaemic) CBF IND pattern in wave 9 to more pronounced hypoperfusion pattern in wave 14, particularly in region of interest b. EG = ectosylvian gyrus.
Figure 5  Secondary infarct growth related to multiple appearance of CSD in a patient suffering from ‘malignant’ ischaemic stroke.  
(A) From the six electrodes of the subdurally implanted electrode strip, ECoG channels A, B, C and D were acquired. CSD associated ECoG depression expanded spatially along the strip electrode from channel D and C to B and finally also A over time. The ECoG signal progressively decreased with repeated CSDs (note the change of scale from 0.7 to 0.2 mV² in channel D after first episode). (B) In addition, the duration of ECoG suppression and ECoG recovery after each CSD was increasingly prolonged indicating a progressive metabolic deterioration in underlying tissue. (C) Repetitive unidirectional propagations of CSDs at intervals with comparable duration in the ischaemic boundary zone (see arrows) appear as slow potential changes of the integrated raw ECoG (upper panel) and as successive reductions of the high-pass filtered ECoG (lower panel). Since each channel displays the potential difference between its two active electrodes, the spread of the slow potential change from one electrode to the next is seen from the phase reversal between two neighbouring channels sharing a common electrode. (D) MRI was conducted after start of monitoring on Day 2 and at the end of monitoring on Day 7 after stroke. The follow-up MRI showed a growth of infarction with secondary ischaemia of the peri-infarct tissue in which 92 CSDs had occurred over the course of around 5 days of monitoring. Note the newly established lesions around the primary infarct area in the diffusion weighted imaging (DWI) on Day 7 corresponding to reduced apparent diffusion coefficients (ADC) in the same regions (see arrow heads) indicating the process of secondary deterioration. The strip electrode (schematically displayed) was placed over peri-infarct tissue, tangentially to the infarct border, with electrodes generating channel D closest to and electrodes generating channel A most distant to the infarct. Note the concentric pattern of cortical infarct growth.
The principal finding in this study was the detection of a wave of cortical blood flow alteration secondary to a depolarization that cycled, spontaneously and several times, around a focal ischaemic lesion in the lissencephalic brain of rats. This gave rise to the observation, at any single given point in the wave path, of recurrent blood flow transients with quite regular periodicity—in temporal terms a ‘cluster’—and the imaging data allowed not only interpretation of such clustering as indicating repetitive peri-lesion cycling of a single depolarization wave, but also demonstrated considerable stepwise enlargement of the ischaemic lesion with each single wave cycle. In the gyrencephalic brain of cats, we identified similar repetitive circumferential propagation around larger ischaemic lesions imaged, however only in a sector of perhaps some 75° of the full 360° of the evolving infarct. Furthermore, the observation that multiple, partially clustered, repetitive PIDs in a patient with malignant hemispheric stroke coincided with major, essentially concentric enlargement of the ischaemic lesion over a period of 5 days led to the inference that the experimental results may apply at least to some extent to human stroke.

The present results demonstrate behaviour similar (in respect of cycling around an anatomical lesion) to that underlying a cardiac arrhythmia that is common in humans, atrial flutter (Murgatroyd and Krahn, 2002), and also indicate that repetitive peri-lesion cycling of a depolarization is a property of cerebral cortex undergoing a naturally occurring pathological process—occlusive ischaemia. Thus peri-lesion cycling does not require surgical lesioning of the cortex to impose a pathway of cyclical propagation, as first described by Shibata and Bures (1972), and is not confined to specially designed preparations of central nervous tissue such as the isolated chick retina (Martins-Ferreira et al., 1974). The findings in rat focal ischaemia accord well with—and offer an explanation for—periodicity in clusters of spreading depolarizations as observed in peri-infarct tissue in the gyrencephalic brain of cats (Ohta et al., 1997) and also in patients with acute ischaemic brain injury (Dreier et al., 2006; Dohmen et al., 2008).

It is remarkable that the circling pattern of PID propagation was consistently observed in rats despite the fact that the experiments in this species were carried out under isoflurane/nitric oxide anaesthesia. This type of anaesthesia is known to attenuate CSD susceptibility in non-ischaemic rat brain (Kudo et al., 2008). The penumbra seems to be more susceptible to CSD than non-ischaemic cortex, based on the observation that known inhibitors of CSD (e.g. NMDA receptor blockers) are less likely to inhibit PIDs in ischaemic penumbral cortex (Shin et al., 2006) or when the cortex is conditioned by artificially elevated extracellular K⁺ (Petzold et al., 2005). Our results indicate therefore that the probability for PID induction and propagation is very high in the penumbra, and might even be augmented under other anaesthetics like 2-chloralose, or in the awake state.

After the very acute phase of ischaemic and metabolic failure and damage, lesion growth into the penumbra may persist for hours or even days (Heiss et al., 1994). Lesion growth ‘pari passu’ with occurrence of individual CSDs has been shown with nicotinamide adenine dinucleotide fluorescence imaging (Strong et al., 1996) and, now in a sequence of CSDs, with LSF (Shin et al., 2006). The present findings are important in showing that sequences of depolarizations can occur and serve to deepen and widen the ischaemia on the basis of repetitive cycling and coupled stepwise deterioration in perfusion around a focus of sustained depolarization in the cerebral cortex—the initial core lesion. [Occurrence of spreading depolarizations in deep grey nuclei is also well documented, both in the human (Sramka et al., 1977) and in the cat (with evidence suggesting temporal clustering) (Umegaki et al., 2005).]

Our review of spatial patterns of spread of depolarizations in the cat MCA occlusion model yielded data compatible with, and strongly suggestive of, cycling of depolarizations around the ischaemic core. Thus we saw examples of circumferential and radial spread, and of depolarizations that split from an origin at the edge of the core, to spread both clockwise and anti-clockwise around the core. Conversely, we recorded more than one instance where depolarizations emerged from either side of the field and collided, suggesting an origin from a single start point diametrically
opposite on the lesion perimeter. However, the limitations imposed by the restricted field of view in this model are significant; for example, dual, simultaneous converging events may alternatively have arisen from new, separate initiation sites at the rim of the core, both of them precipitated by a transient reduction in arterial pressure. Given the restricted fraction of the ischaemic territory that can be imaged in these experiments, greater reliance has to be placed on the demonstration of recurrent depolarizations at reasonably repetitive time intervals (perhaps within 10–15%) as evidence for cycling. That the incidence of tight periodicity is not as high in the cat MCA occlusion preparation as with rat dMCAO does not argue against this hypothesis; this is because the longer perimeter of the core in the cat brain raises the probability of additional, new depolarizations originating at the perimeter and interrupting regular repetition of a cycling wave.

Are we justified in applying the spatial cycling model to human ischaemic brain injury? The relatively tight periodicity of depolarizations shown in Fig. 5 from a patient with malignant hemispheric stroke supports this suggestion. However, although the increase in modal interval between clusters of depolarizations as we move from rat to cat brain appears appropriate in relation to the likely absolute lesion perimeter, this argument cannot be sustained when comparing intervals between events in cat versus human brain. A much longer modal interval would be expected of cycling around an MCA infarct in humans than is indicated in Fig. 6. The discrepancy may be explained in two ways (that are not mutually exclusive). First, the much longer (absolute) lesion perimeter in humans (MCA occlusion) increases the probability of new events arising at the perimeter, interrupting any regular cycling observed at the relatively localized site of a single recording strip on the lesion periphery. Second, where clear periodicity is seen with a modal interval of the order we describe, this would suggest the possibility of cycling around a smaller, evolving ‘satellite’ core situated within what is still largely penumbra. Such heterogeneity has been described, at least in experimental stroke models (Tomlinson et al., 1993).

Critically, the demonstration of repetitive peri-lesion cycling of a depolarization introduces the novel hypothesis that this behaviour may not simply cause lesion growth but also serve as a biological amplifier, enhancing upregulation around a lesion of inflammatory, stress- and neurogenic-response cascades to focal brain injury that are known to be related to spreading depolarization (Sharp et al., 2000; Jander et al., 2001; Gursoy-Ozdemir et al., 2004; Yanamoto et al., 2005). This could potentially apply to any tissue in the central nervous system capable of supporting CSD, namely a concentration of neuronal cell bodies, astrocytes (or Müller cells in the retina) and their associated vasculature—whether in the cerebral cortex, deep nuclei (Sramka et al., 1977), brainstem (Richter et al., 2003), spinal cord (Czeh and Somjen, 1990) or retina. As an example of a pathogenic effect of CSDs, matrix-metalloproteinase-9 upregulation increases blood–brain barrier permeability (Gursoy-Ozdemir et al., 2004) and hence can promote vasogenic oedema. On the positive side, deliberate preconditioning of the rat cerebral cortex with CSD confers a measure of protection against a subsequent ischaemic challenge (Kobayashi et al., 1995). However, it is very clear now that repetitive CSDs under conditions of focal ischaemia—PIDs—cause vasoconstriction in the microcirculation (Dreier et al., 1998; Shin et al., 2006; Strong et al., 2007) and hence promote lesion expansion and/or deterioration towards terminal depolarization and infarction. The data in Fig. 1C (circumferential spread of depolarization: rat) and Supplementary Video 2 (nakamura_suppl_v2.mov) (radial spread: cat) suggest that the time course of the perfusion response to PID is biphasic nearest the core (hyperaemia followed by hypoperfusion in the rat: hypoperfusion followed by mild hyperaemia in the cat) but monophasic (hyperaemia: either species) more remotely and nearer sources of collateral perfusion. This is consistent with the observation of impaired neurovascular coupling at the infarct border (Kunz et al., 2007), although the relative phasing of hyperaemia and hypoperfusion may vary between species (Strong et al., 2007). The precise features of ischaemia that determine the apparently critical reversal of the vascular response to depolarization from dilation to constriction remain to be identified, but recent work suggests that downregulation of the endothelial nitric oxide synthase pathway contributes to this reversal (Petzold et al., 2008).

The present results do not by themselves prove that PIDs are pathogenic in the human brain, in terms of specific histological or structural MRI findings that can be related to number and/or patterns of depolarizations. A prospective study of this nature is desirable. However, three elegant experimental studies suggest that number of PIDs is an independent predictor of infarct size (Back et al., 1996; Busch et al., 1996; Takano et al., 1996). In addition, new clinical monitoring data indicate that some depolarizations may be accompanied by reductions in cerebral blood flow suggesting vasoconstriction (Dreier et al., 2009), and often by depletion of the interstitial brain glucose pool (Feuerstein et al., 2010), suggesting a pathogenic potential.

These features, especially when set against the suggestion (above) that CSD associated with the (more normal) hyperaemic response may be protective, raise the issue of clinical management. Briefly, we submit that clearer evidence of the relationships of different patterns of depolarizations—dilator or constrictor—with outcome is needed before specific therapy can be designed. Clinical application would probably be dependent first on distinction of these two patterns in individual patients, and should recognize (i) that blockade of hyperaemic (normal response) CSD may compromise a response that is potentially protective and (ii) that PIDs are unlikely to respond to glutamate antagonists (Shin et al., 2006) [although earlier experimental work with a gyrencephalic MCA occlusion stroke model suggested that post-treatment with MK801 is protective (Park et al., 1988), and there is initial clinical evidence to suggest that it may sometimes be possible to block depolarizations (Sakowitz et al., 2009)]. Until it is clear which patterns of depolarization are independently associated with better or worse outcome, medical management should probably focus simply on careful application of some principles of good clinical management: avoidance of hypotension and pyrexia (Hartings et al., 2009) and maintenance of an adequate but not excessive plasma glucose, most probably in the range 7–9 mmol/l (125–160 mg/100 ml) (Oddo et al., 2008; Strong, 2008).

In conclusion, the present findings explain observations of periodicity in CSD occurrence in acute brain injury and the impact of
CSDs on infarct size. Future work should address what determines (i) variation between adverse and restorative-protective responses and (ii) clinical outcome in human brain injury, and only then should focus on the central goal of effective therapy.

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Supplementary material

Supplementary material is available at Brain online.

References


