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ORIGINAL ARTICLE

Monitoring stroke progression: *in vivo* imaging of cortical perfusion, blood–brain barrier permeability and cellular damage in the rat photothrombosis model

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Focal cerebral ischemia is among the main causes of death and disability worldwide. The ischemic core often progresses, invading the peri-ischemic brain; however, assessing the propensity of the peri-ischemic brain to undergo secondary damage, understanding the underlying mechanisms, and adjusting treatment accordingly remain clinically unmet challenges. A significant hallmark of the peri-ischemic brain is dysfunction of the blood-brain barrier (BBB), yet the role of disturbed vascular permeability in stroke progression is unclear. Here we describe a longitudinal *in vivo* fluorescence imaging approach for the evaluation of cortical perfusion, BBB dysfunction, free radical formation and cellular injury using the photothrombosis vascular occlusion model in male Sprague Dawley rats. Blood-brain barrier dysfunction propagated within the peri-ischemic brain in the first hours after photothrombosis and was associated with free radical formation and cellular injury. Inhibiting free radical signaling significantly reduced progressive cellular damage after photothrombosis, with no significant effect on blood flow and BBB permeability. Our approach allows a dynamic follow-up of cellular events and their response to therapeutics in the acutely injured cerebral cortex.

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INTRODUCTION

Focal ischemic stroke is among the main causes of death and morbidity worldwide.¹ Thrombolytic therapy remains the only approved treatment for acute ischemic stroke, yet it only benefits a limited number of patients diagnosed and treated within the first few hours after the insult.^{2,3} Patients' outcome was shown to be critically determined by alterations in the brain tissue surrounding the ischemic core; for example, in the vicinity of the lesion core, nonischemic brain regions are often characterized by reduced blood flow and disturbed neural functions.⁴ Recent studies pointed to vascular dysfunction, specifically impairment in blood-brain barrier (BBB) permeability as an additional important hallmark of the peri-ischemic brain.⁵ Indeed, changes directly related to BBB opening seem to play a critical role in the pathogenic process after stroke. Kuntz et al⁶ recently showed BBB dysfunction coinciding with cell damage in a cell culture model of ischemia and reperfusion, and after transient medial cerebral artery occlusion in mice. Experimental BBB injury in rodents leads to neuronal dysfunction, spreading depolarization, seizures and delayed degeneration-all well-known complications of stroke.^{7–11} In stroke patients, BBB dysfunction is associated with increased risk of secondary hemorrhage and cerebral edema,

further compromising perfusion because of vasospasms or through increased intracranial pressure.^{2,7,12,13} However, the spatiotemporal dynamics and mechanisms underlying vascular alterations and the potential link between BBB opening, neurovascular dysfunction and cellular injury after stroke are not fully understood.

We studied photothrombosis (PT)-induced changes in cortical perfusion, BBB permeability, free radical formation and cell injury in vivo-focusing on the longitudinal experimental design to allow a parallel study of the spatiotemporal dynamics of these parameters. Cortical perfusion and BBB permeability were quantified using fluorescent angiography by intravenous application of fluorescein sodium salt, commonly used in the clinics to assess retinal blood flow and blood-retinal barrier permeability;¹⁴ in these studies the first pass of the tracer bolus (representing vascular filling) is used to determine tissue perfusion and the delayed extravascular (EV) accumulation of the tracer reflects its leakage from injured vessels.^{15–17} Formation of reactive oxygen and nitrogen species were monitored using different fluorescent probes. Nitric oxide (NO) was detected using 4-amino-5-methylamino-2',7'-difluorofluorescein diacetate (Molecular Probes, Eugene, OR, USA; its oxidative derivatives being reactive with NO) as

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previously used in a transient medial cerebral artery occlusion model in mice,¹⁸ and reactive oxygen species (superoxide and hydroxyl radicals) were detected by ROSstar 650 (Li-Cor, Lincoln, NE, USA), a hydrocyanine-based indicator, previously used in vivo after lipopolysaccharide-induced peritoneal inflammation and implant-associated inflammation.^{19,20} ROSstar 650 (Li-Cor) is membrane permeable in its reduced form and becomes fluorescent and membrane impermeable once oxidized, hence can be used for the detection of intracellular reactive oxygen species (ROS) formation.¹⁹ Cellular injury was detected using propidium iodide (PI), a membrane integrity marker that binds to DNA/RNA and increases its fluorescence by a factor of 20 to 30 upon binding, and fluorescently labeled annexin V, an endogenous protein with high affinity to phosphatidylserine. Phosphatidylserine becomes exposed to the outer leaflet of the membrane during apoptosis and necrosis and has been successfully detected in animal models of cerebral ischemia and in a human study.^{21,22} As ROS were shown to be involved in deleterious processes within the periischemic brain,²³⁻²⁵ we aimed to use our imaging approach to follow the spatial and temporal progression of ROS synthesis after the insult. This approach enabled us to test the modifying effect of inhibiting free radical signaling on the dynamics of cortical perfusion, BBB permeability and cell damage in the periischemic brain.

MATERIALS AND METHODS

All experimental procedures were performed according to the guidelines of the animal care and ethical committee at Ben-Gurion University of the Negev, Beer-Sheva, Israel.

Photothrombosis Model of Vascular Occlusion

A total of 34 male Sprague Dawley rats (bodyweight 235 to 380 g, Harlan, Jerusalem, Israel), housed under standard conditions with free access to food and water, underwent craniotomy over the right somatosensory cortex (2 mm frontal—4 mm occipital and 2 to 6 mm lateral to bregma) under deep anesthesia (intraperitonal injection of ketamine (100 mg/mL, 0.08 mL/100 g) and xylazine (20 mg/mL, 0.06 mL/100 g)).¹⁶ Oxygen saturation was continuously monitored at the hind paw (Starr Life Science MouseOx probe, Oakmont, PA, USA) and body temperature was kept at $37^{\circ}C \pm 0.5^{\circ}C$ using a heating pad. After removal of the dura, Rose bengal (RB, Sigma-Aldrich, St Louis, MO, USA) was injected into the tail vein (0.133 mL/ 100 g bodyweight, 7.5 mg/mL saline, 9.8 μ mol/l/kg) and PT was induced using focal illumination (532 nm, diameter of ~ 1 mm, CNI Laser, MGL-III-532-5 mW-1.5, Changchun, China) for 15 minutes.²⁶ The animal was strictly shielded from light, minimizing diffuse RB activation after laser illumination.

Intravital Fluorescence Imaging

The imaging-setup consisted of a fluorescence microscope (Zeiss SteReO Lumar V12; Oberkochen, Germany) with three built-in filters and an EMCCD camera (Andor Technology, DL-658 M-TIL; Belfast, UK). Images were captured at x6.4 to x40 magnification with quantitative analysis based on x30 magnification and a region of interest of 2.19 mm × 1.65 mm. Perfusion and BBB permeability assessment were based on intravenous injections of fluorescein sodium salt (1 mg/kg bodyweight, 1 mg/mL saline, Sigma-Aldrich, St Louis, MO, USA) pre-PT and hourly after PT. Images were taken at 5 Hz for 306 seconds starting 6 seconds before injection. To detect cell death/damage, recombinant human annexin A5 (from Chris P Reutelingsperger) was labeled with fluorescein isothiocyanate (FITC, Sigma-Aldrich, St Louis, MO, USA) as described.²⁷ Two forms were used: annexin A5 with retained affinity for phosphatidylserine and nonbinding annexin A5 with a complete loss of affinity for phosphatidylserine. Both FITC-labeled annexin A5 preparations were diluted in saline from FITC annexin stock (nonbinding: 1 g/l protein, cFITC 22.2 µmol/l, FITC/ annexin ratio 0.8; binding: 1.2 g/l protein, cFITC 20.5 µmol/l, FITC/annexin ratio 0.6) and ~25 nmol of FITC/100 g bodyweight were injected intravenously into the tail vein before and after PT (n = 4). Propidium iodide (Molecular Probes, Eugene, OR, USA; 0.5 mg/kg bodyweight, 0.5 mg/ml saline) was intravenously injected before and hourly after PT or superfused topically. Topical application of PI, to bypass the BBB, did not prove feasible as widespread surface noise was always detected (n = 4, data not shown). Propidium iodide and FITC-annexin A5 signals were acquired before tracer injection and 10 minutes after injection at 1 Hz. For quantitative image analysis of PI, an averaged image (consisting of 10 frames) was created.

ROSstar 650 (Li-Cor) to detect superoxide and hydroxyl radicals was applied by intravenous injection (0.2 to 0.5 mL saline-based solution, concentration: 50 to 100 μ mol/l) 30 minutes before the induction of PT and reinjected up to 90 minutes after RB injection (n = 2) or was topically applied and incubated for 30 minutes before imaging (n = 1). 4-Amino-5-methylamino-2',7'-difluorofluorescein diacetate (Molecular Probes) was topically applied (concentration: 50 to 200 μ mol/l, 30 minutes incubation)¹⁸ before PT for baseline imaging and ~ 3 hours after PT to detect NO (n = 5).

For BBB imaging in deep cortical layers *ex vivo*, 0.2 mL/ 100 g bodyweight of 2% Evans blue solution (Sigma-Aldrich, St Louis, MO, USA) was injected into the tail vein 1 hour before sacrificing the animal as previously reported.^{8,28,29}

Inhibition of Reactive Oxygen Species

To inhibit reactive oxygen/nitrogen species and their generation, a combination of 1 mmol/l 2-(4-carboxyphenyl)-4,4,5,5-tetramethyl-imidazoline-loxyl-3-oxide (carboxy-PTIO, NO scavenger; Tocris, Bristol, UK), 1 mmol/l L-N^G-Nitroarginine methyl ester (NO synthase inhibitor; Tocris, Bristol, UK) and 1 mmol/l phenyl-*N*-t-butylnitrone (free radical spin trap (superoxide) and inhibitor of induction of NO synthase; Sigma-Aldrich, St Louis, MO, USA) were used and perfused in artificial cerebrospinal fluid. Animals were assigned randomly to the treatment group (n = 9). In the nontreated group (n = 9), two animals were excluded because of BBB opening before PT induction. Since the PT model is based on free radical generation (by activated RB), and to ensure that free radical inhibiting treatment will not affect clot generation and development of the ischemic core, we started topical perfusion (0.5 ml/h) with free radical inhibitors 30 minutes after RB injection when >90% of blood RB was hepatically eliminated.³⁰

Ex Vivo Microscopy and Histology

Snap-frozen brains were cut in 20 μ m thick coronal slices with a cryostat (Leica CM 1950, Wetzlar, Germany) and rinsed in an alcohol series of decreasing concentrations before incubation with hemalaun (Merck, Darmstadt, Germany). The slices were then rinsed in 96% ethanol with 2% hydrochloric acid and incubated in 2% sodium bicarbonate. After dehydration and mounting, the sections were analyzed with a LEICA DMRE microscope (Leica Microsystems Jena Gmbh, Jena, Germany). For detection of annexin A5 and PI signals, *ex vivo* nonstained adjacent slices were used.

Data Processing and Analysis

Imaging data were transferred to MATLAB and corrected for movement artifacts as described.¹⁶ To estimate relative perfusion, we first extracted the arterial input function (AIF, dynamic arterial signal intensity after tracer injection) from an arteriole (at least one branching point away from the occluded vessel) that stayed perfused throughout the experiment as documented by repeated fluorescent angiography. The ratio between the maximal intensity change in each pixel during first pass (FP_Max) and the maximal AIF (maxAIF) was termed 'relative perfusion' and taken as a surrogate marker for local cerebral perfusion. The 'lesion border' was defined by post PT angiography, which delineated the ischemic core, where arteriolar supply was blocked, from the peri-ischemic brain. For each pixel relative perfusion was presented as a function of distance from this border. Averaged relative perfusion values were calculated for pixels in eccentric rings of 100 µm width starting at the lesion border and normalized to the control value measured before PT (normalized relative perfusion = (FP Max(PT)/maxAIF(PT))/(FP Max(Pre PT)/maxAIF(Pre PT)) \times 100%). To quantify the area most severely affected by reduced perfusion, we arbitrarily set a perfusion threshold at 30% maxAIF during tracer first pass. Although > 99% of the pixels were above threshold in the intact brain, thus identified as 'normally perfused', the majority of pixels within the ischemic core $(70.0\% \pm 12.0\%)$ were characterized as 'hypoperfused' according to this threshold.

To assess vascular permeability, the EV and vascular compartment were segmented on the basis of an arteriovenous first-pass overlay image and the 'adaptive threshold' function in MATLAB.²⁹ For each EV pixel, the integral of the decay phase of the postinjection curve was computed and divided by the AIF integral (EV/AIF), thereby comparing the change in EV signal intensity (reflecting tracer accumulation) relative to the arterial

input. We refer to this ratio as the 'BBB permeability index'. Cellular damage was quantified as the number of PI-positive pixels after background correction. Both BBB permeability and cell damage were assessed as functions of distance to the lesion border (as described above).

Statistical Analysis

Α

Cranial window

Statistical analysis was performed with nonparametric Friedman's analysis of variance for related samples and nonparametric Mann–Whitney U-test

R

Cntrl

20 min PT

EΒ

Model

for independent samples with Bonferroni *post hoc* correction. Differences were considered significant at P < 0.05. All statistical analyses were performed with SPSS v.20.0 (IBM Corporation, Armonk, NY, USA). Data is shown as mean \pm s.e.m.

RESULTS

Nobind Anx 140 min PT

Nobind Anx FF

Anx A5 150 min PT

PI 160 min PT

PI/ Anx A5 OL

Cellular damage in vivo

Photothrombosis was induced through simultaneous RB injection and laser illumination, impairing blood flow within seconds

Histology

Anx A5

PI

PI/ Anx A5 OL



С



(see Supplementary Movie). Consistent with previous studies, macroscopic sections confirmed BBB breakdown in most layers of the cortex surrounding the lesion core (Figure 1A, bottom).^{8,28,29} Cellular damage was detected in the peri-ischemic region using intravital microscopy after intravenous injection of either PI or binding annexin A5, while nonbinding annexin A5 gave no specific staining (Figure 1B). Positive annexin A5 staining, while generally colocalized with PI (n = 4), was most prominent in close proximity to blood vessels, mainly venules. Notably, in the periphery of the peri-ischemic regions, positive annexin A5 staining was not always associated with PI fluorescence (Figure 1B, arrows). Similar to BBB dysfunction (Figure 1A, bottom), ex vivo microscopy confirmed cellular injury in most cortical layers (Figure 1C). For quantitative analysis of cell injury we used PI images, since PI gave a better signal to noise ratio compared with annexin, perhaps because of its small molecular size and rapid diffusion or because of increased fluorescence upon DNA/RNA binding.³¹

Using repeated angiography, we quantified dynamic changes in relative perfusion within each treated brain, allowing the characterization of three regions: (1) an 'ischemic core'—where PT was induced and arterial supply permanently blocked; (2) an *adjacent peri-ischemic zone* with significantly reduced blood flow compared with the pre-PT state (< 400 μ m); and (3) a *distant peri-ischemic zone*—in which perfusion was intact (similar to pre-PT state) throughout the 3 hour monitoring period (Figure 2).

Longitudinal *in vivo* imaging showed that reduction in perfusion, increased vascular permeability, and cellular damage clearly progressed within the peri-ischemic brain after PT (Figures 2C–F, Figure 3). Notably, while reduced blood flow was limited to the adjacent peri-ischemic zone, BBB permeability and cell damage (PI signal) were increased within both the adjacent and distant peri-ischemic region (Figures 3B–D, see quantification below and Figure 6).

To study the involvement of free radical signaling in the peri-ischemic brain, we monitored the formation of NO (using 4-amino-5-methylamino-2',7'-difluorofluorescein (DAF-FM) diacetate), and of superoxide and hydroxyl radicals (using ROSstar 650 (Li-Cor)) in vivo. Oxidized ROSstar 650 (Li-Cor) and the NOadduct of DAF-FM diacetate were detected in the ischemic core and within the peri-ischemic cortex (Figures 4 and 5). Although PT was followed by a diffuse vascular and parenchymal signal of ROSstar 650 (Li-Cor; n = 3; Figure 4), NO staining delineated arterioles (n = 5, Figures 5C and 5D). The delayed detection of free radicals in the peri-ischemic brain suggests that free radical formation was independent of direct RB activation. We further investigated whether inhibition of ROS signaling affected cerebral perfusion, BBB permeability, and cell damage after PT by applying a 'cocktail' containing phenyl-N-t-butylnitrone (free radical spin trap), L-N^G-nitroarginine methyl ester (NO synthetase inhibitor), and carboxy-PTIO (NO scavenger) 30 minutes after PT. We applied this cocktail, rather than selective blockers, because reactive oxygen as well as nitrogen species (especially peroxynitrite) were shown to be involved in the no-reflow phenomenon after medial cerebral artery occlusion,²⁵ and in cellular damage.^{23,24}

Quantitative analysis of relative perfusion, BBB permeability, and cellular injury with and without ROS inhibition (treatment: n = 9; no treatment: n = 7) confirmed lesion progression in the PT model. Without treatment, $11.3\% \pm 2.9\%$ of the adjacent zone was hypoperfused (below 30% maximal AIF, P = 0.04) 1 hour after clot formation, and $25.1\% \pm 4.4\%$ was hypoperfused after 3 hours (Figure 6A, P < 0.001). No significant perfusion deficit was observed in the distant peri-ischemic zone. Reactive oxygen species inhibition reduced the percentage of hypoperfused tissue in the adjacent zone to $4.8\% \pm 1.4\%$ and $13.2\% \pm 3.1\%$ at 1 and 3 hours after PT, respectively, however not reaching significance (P = 0.102 and 0.041, respectively before *post hoc* correction; Figure 6A).

Vascular permeability to the injected sodium fluorescein was significantly increased in the adjacent zone one hour after PT ($11.11\% \pm 1.88\%$, P < 0.001) and propagated to the distant perischemic region (2.9 ± 1.76 , P = 0.01) at 3 hours. ROS inhibition had no effect on the progression of BBB permeability changes (Figure 6B).

Quantitative analysis of the PI signal confirmed propagation of PT-induced cellular damage within the peri-ischemic brain (Figure 6C). The number of PI-positive pixels significantly increased within the adjacent peri-ischemic region when observed 1 hour after PT ($2.7\% \pm 1.1\%$ versus $0.002\% \pm 0.002\%$, post- versus pre-PT, respectively, P = 0.003). After 3 hours, PI signal was significantly increased in both the adjacent ($17.2\% \pm 2.6\%$ PIpositive pixels, P < 0.001) and distant zone ($4.9\% \pm 1.6\%$ versus $0.9\% \pm 0.5\%$ PI-positive pixels, post- versus pre-PT, P = 0.002; Figure 6C). Although cellular injury in the adjacent zone remained unaffected by ROS inhibition, the treatment effectively prevented the progression of cellular damage to the distant peri-ischemic region, where cellular damage was significantly reduced to $0.3\% \pm 0.2\%$ of PI-positive pixels (P < 0.001, Figure 6C).

DISCUSSION

Using longitudinal *in vivo* imaging within individual animals, we monitored the progressive nature of lesion development after clot formation. During the first 3 hours after PT we show (1) a progressive reduction in blood flow, BBB dysfunction, and cellular damage in the peri-ischemic brain; (2) progression of BBB breakdown and cell damage exceeding the hypoperfused brain areas; (3) synthesis of reactive oxygen species within the lesion core and its surroundings; and (4) reduced progression of cell damage into the distant perilesional zone under inhibition of free radical signaling, without significant effects on changes in perfusion and permeability.

The classic stroke literature describes an ischemic core and a surrounding brain region of dysfunctional brain parenchyma with reduced perfusion that may escape cell death (referred to as 'benign oligemia') or may undergo delayed infarction (true 'tissueat-risk').^{4,32} However, defining thresholds and measuring levels of cerebral perfusion to distinguish regions with 'benign oligemia' from true 'tissue-at-risk' remains challenging. This is not only because of the limitations of available imaging modalities (mainly computed tomography and magnetic resonance imaging

Figure 2. Partitioning the peri-ischemic brain according to perfusion measurements. (**A**) Schematic of the ischemic core (IC), the adjacent (AZ), and distant peri-ischemic zone (DZ, lesion border, LB). (**B**) Flow curves, taken from the fluorescent angiography illustrate reduced perfusion in AZ and IC, while the reference arteriole (AIF, arterial input function) and DZ remained unaffected. PT, photothrombosis. (**C**) Relative perfusion maps by pixel-based normalization of the first pass maximum (FP_Max, see B) to the maximal AIF (maxAIF). (**D**) Relative perfusion averaged by distance to LB. (**E**) Relative perfusion normalized to pre-PT perfusion calculated as follows: ##normalized relative perfusion = (FP_Max(PT)/maxAIF (PT))/(FP_Max(Pre PT)/maxAIF(Pre PT)) × 100%. (**F**) Quantitative analysis reveals that reduced normalized relative perfusion remains within 400 μ m from LB during the 3 hours monitoring period (n = 7) allowing the division of the peri-ischemic brain into AZ and DZ. For statistical analysis nonparametric paired Friedman's analysis of variance including Bonferroni *post hoc* correction was performed; adjusted *P*-values in control comparison ***P < 0.001; **P < 0.01; *P < 0.05. Data shown as mean \pm s.e.m. *For graphical simplicity only largest distance shown on axis label (e.g., '200' contains mean of all values between 100 and 200 μ m); scale bar = 200 μ m.

perfusion), e.g., utilizing nonuniform image processing,³³ but also because of the dynamic nature of stroke pathophysiology itself resulting in varying viability thresholds at different time points

after stroke.³² In our study, regional differences in cerebral perfusion were measured using fluorescent angiography, previously shown to correlate with laser Doppler flowmetry.^{16,29} Angiography



Journal of Cerebral Blood Flow & Metabolism (2014), 1791 – 1801

1795



Figure 3. Parallel intravital monitoring of relative cerebral perfusion, blood-brain barrier (BBB) permeability, and cell damage. (**A**–**C**) From top to bottom: images under control condition and 1 to 3 hours after Rose bengal-induced photothrombosis. (**A**) Arteriovenous first pass images. Pixels were counted as hypoperfused when below a threshold of 30% of the maximum of the arterial input function (AIF—selected arteriole marked by asterisk and red dotted line). The white dotted line marks the hypoperfused lesion core. (**B**) BBB permeability maps. Dotted square marks region of interest shown in C. Dotted lines were transferred from A. (**C**) Cell damage as detected by propidium iodide (PI). Note how cell damage progresses into normally perfused perilesional brain tissue. Increases in BBB permeability seem to precede cell damage. (**D**) The graphs show how hypoperfusion, increased BBB permeability, and cell damage affected growing perilesional areas with time after PT. AZ, adjacent zone; DZ, distant zone. Scale bar = 200 μ m.



В

Superoxide/ Hydroxyl-Radical (ROSstar 650)



Figure 4. Intravital microscopy reveals superoxide and hydroxyl radical formation after photothrombosis (PT). (**A**) Left: bright field image 40 minutes after PT induction. Right: blood-brain barrier (BBB) permeability index 80 minutes after PT. (**B**) Areas with enhanced superoxide and hydroxyl radical formation. Note the detection of free radicals in the ischemic core as well as in the peri-ischemic cortex. (**A** and **B**) Dotted lines indicate previously laser-illuminated ischemic core. Scale bar = $200 \,\mu$ m.

allows measuring blood flow in a higher spatial, yet limited temporal resolution compared with laser Doppler flowmetry. Most importantly, using this technique, we were able to quantify changes in vascular permeability owing to leakage of tracer through the dysfunctional BBB.^{16,34} We describe an ischemic core (~1.2 mm diameter) surrounded by a rim with a width of ~ 400 μ m that was characterized by progressive hypoperfusion and cell damage during the 3 hours of monitoring. This region is consistent with the small peri-ischemic hypoperfused rim described 1 to 6 hours after stroke in patients, that may predict tissue deterioration.³⁵

We used intravenous annexin A5 and PI to follow cellular damage *in vivo*, as these were previously shown to increase in models of brain injury and confirmed by terminal transferasemediated dUTP nick-end labeling costaining *ex vivo*.^{21,22,36} Although PI, as a marker for membrane integrity, is assumed to detect irreversible damage, phosphatidylserine exposure was shown to be reversible under mild stress. Hence annexin A5-positive cells may potentially escape cell death.³⁷ We saw a large overlap between PI and annexin A5 staining in the peri-ischemic brain. In an earlier study using the PT model, the largest fraction of

apoptotic cells surrounding the infarct core were morphologically identified as neurons.³⁸ Similarly, Van Hoecke *et al*³⁹ describe a 'surrounding pale region' around the ischemic core characterized by significant neuronal cell death forming within the first 12 to 24 hours after PT. Conversely, resident microglia, blood-borne leukocytes and astrocytes start to align and proliferate around the lesion core within the first 24 to 48 hours.^{40,41} Interestingly, within the periphery of the peri-ischemic zone we detected annexin A5-positive, PI-negative cells, suggesting reversible cellular injury (see Figure 1B, arrowheads). Similar results showing peripheral and contralateral annexin A5-positive cells have been published in a transient medial cerebral artery occlusion model.²² An alternative hypothesis explaining annexin A5-positive regions without Pl-uptake in our experiments would be a difference in vessels' permeability, selectively allowing the passage of large-molecularweight proteins, such as annexin A5 protein, but not the smaller, yet charged, PI (BBB nonpenetrable),36 perhaps because of modulation of transcellular and not paracellular vascular barriers.⁴² However, the latter hypothesis, which may have led to underestimation of cell damage in our experiments utilizing PI, is







Figure 5. Intravital microscopy reveals nitric oxide (NO) formation after photothrombosis (PT). (**A**) From left to right: bright field images under control condition and 80 and 160 minutes after induction of PT. Note the lesion core in the top left corner. (**B**) Cell damage as visualized by propidium iodide (PI). (**C**) Nitric oxide as detected by 4-amino-5-methylamino-2',7'-difluorofluorescein (DAF-FM) diacetate gave a diffuse parenchymal signal before PT and clearly showed arteriolar NO production at 80 and 200 minutes after PT in the ischemic core (here veins also revealed signal) and in the peri-ischemic arterioles. (**D**) Overlay of PI and DAF-FM. Note how cell damage is mainly concentrated surrounding venules. (**A**, **C**, and **D**) asterisk indicates arteriole that was primarily perfused after PT, however, underwent delayed constriction despite NO production. Scale bar = 200 μ m.

less likely, since the region displaying BBB dysfunction (leakage of the charged tracer sodium fluorescein) clearly exceeded the region affected by cell damage, especially in animals treated with ROS inhibitors (Figures 3 and 6).

Our observation of progressive cell damage in the distant, apparently normally perfused, brain tissue is consistent with the notion that perfusion-based imaging methods may not detect regions eventually showing mild cell damage, and with the concept of selective neuronal loss in remote peri-ischemic regions.⁴³ Interestingly, we show here that normally perfused, remote peri-ischemic regions, are characterized by BBB dysfunction together with cellular damage. Accumulating results thus highlight imaging methods for the detection of a compromised BBB, as potentially sensitive for the detection of perilesional tissue at risk. In the nonischemic brain, experimental induction of BBB opening has been shown to induce neuronal damage preceded

С

Nitric oxide (DAF-FM)

D

DAF-PI Overlav (B+C)

Imaging experimental stroke progression *in vivo* K Schoknecht *et al*



Figure 6. Quantification of perfusion deficit, blood-brain barrier (BBB) permeability, and cell damage after photothrombosis—involvement of free radical signaling. (**A**) Perfusion deficit was assessed as percentage of area not crossing 30% of the maximum of the arterial input function (maxAIF). Although under control conditions < 1% of the area was subthreshold ('hypoperfused', combined groups PT (n = 7) and PT+treat (n = 9)), the hypoperfused area increased in both the treated and nontreated group. Treatment reduced the hypoperfused area, however, not reaching significance. The distant peri-ischemic zone (400 to 800 μ m from ischemic core) remained largely unaffected. (**B**) Blood-brain barrier permeability increased early and robustly in the adjacent peri-ischemic zone with progression to the distant peri-ischemic zone 3 hours after photothrombosis. Blood-brain barrier alterations remained unaffected by treatment. (**C**) Mild increases in cell damage were observed early in the adjacent peri-ischemic zone was reduced by inhibition of free radical signaling. Statistical analysis was performed with nonparametric paired distant peri-ischemic for related samples and Mann–Whitney *U*-test for independent samples including Bonferroni *post hoc* correction. Each ring of 100 μ m width was represented by one mean value. ***P < 0.001; **P < 0.05. AZ, adjacent zone; DZ, distant peri-ischemic zone; PT, photothrombosis.

by the activation of astrocytes through serum-derived albumin and activation of inflammatory cascades.^{6,9,11,44} Similar pathologic findings (e.g., glial activation) have been described in studies on selective neuronal loss and in the peri-ischemic brain after PT.^{29,43} Future studies are awaited to confirm the role of BBB damage in nonischemic cellular damage and offer therapeutic strategies to facilitate its repair.

Several mechanisms have been described to underlie hypoperfusion, BBB dysfunction, and cell damage, among them energy depletion, excessive glutamate release, ionic imbalance (e.g., calcium overload, increased extracellular potassium), ROS production, peri-infarct depolarization and inflammatory response (e.g., via cytotoxic cytokines). Although morphologic correlates of the cellular immune response such as activated microglia, astrocytes, and blood-borne macrophages are only present ~1 day after the stroke, earlier deleterious events have been suggested to converge to free radical formation including excessive glutamate release, calcium overload, energy depletion, protease activation, NO synthetase uncoupling, and mitochondrial depolarization.^{24,25,45} Conversely, NO may be essential to maintain or increase collateral blood flow in the peri-ischemic brain (see below) and reactive oxygen and nitrogen species were shown to be relevant cellular signaling molecules, involved in modulation of protein function and in angiogenesis.^{46,47} Hence, prolonged interference with free radical signaling could be deleterious as it affects physiologic functions—a point that was raised to explain why clinical trials using free radical scavengers after stroke have largely failed.4

In vivo detection of free radical formation remains technically challenging. Despite the recent development of new small molecule indicators, most data still originate from *in vitro* models.⁴⁸ Recently, DAF-FM was successfully used to detect NO formation in the arterial and venous cerebral circulation after transient medial cerebral artery occlusion in the mice brain *in vivo*,¹⁸ yet it is still to be uncovered which cells produce NO.²⁴ We report increased NO levels in the peri-ischemic brain, mostly in close vicinity to the arterial wall, suggesting a major nonneuronal contribution to NO formation. The source of superoxide and hydroxyl radicals remains elusive as we detected both vascular and parenchymal signal.

Nitric oxide and superoxide together form toxic peroxynitrite that mediates pericyte constriction,²⁵ which in turn may lead to progressive cellular injury. The fact that we inhibited both superoxide/ hydroxyl and NO formation, hence inhibited the NO scavenging effect of superoxide (which would result in peroxynitrite formation), may explain the lack of vasoconstriction despite NO scavenging. Although in the Yemisci et al²⁵ study inhibition of free radicals was associated with reversed constriction, we found a moderate (yet insignificant) reduction in hypoperfusion after free radical inhibition-perhaps because of topical (not intravenous) application in our study, the limited monitoring period (3 hours after PT in contrast to ex vivo analysis 6 hours after ischemia by Yemisci et al) or the small sample size. Our data are also consistent with the report by Kuntz et al⁶ showing that BBB leakage coincides with emerging cell death. However, it is important to note that reducing cellular injury via free radical inhibition did not affect the extent of BBB breakdown, suggesting that BBB dysfunction in the distant peri-ischemic zone is not due to cell damage. The limited effect of ROS inhibition on perfusion is in line with recent studies in the slice preparation showing ischemia-induced pericyteconstriction associated with pericytic death shortly thereafter, and no significant involvement of ROS.⁴⁹ A surprising result was the lack of effect of ROS inhibition on the progression of BBB damage. Free radicals were previously shown to underlie ischemia-induced BBB dysfunction,⁵⁰ however, a major difference between the studies is the absence of reperfusion in the PT model we used. Reperfusion is known to result in a burst of additional ROS formation,⁵¹ hence may aggravate ROS-induced neurovascular damage. Mechanisms underlying BBB breakdown may differ between models and brain regions exposed to permanent ischemia-and occlusion/reperfusion. Alternatively, since we started free radical inhibition 30 minutes after PT, there is still a theoretical possibility that free radical formation within that time window, initiated a cascade of events (even in the distant region) that lead to delayed BBB dysfunction.

In conclusion, the strength of our study lies in the parallel assessment of different physiologic parameters *in vivo*, allowing further in-depth studies into the mechanisms underlying brain damage in stroke models and testing of new therapeutics. The

nature of the PT lesion (reproducibility, clear ischemic core, and peri-ischemic tissue visible at the brain surface) is ideal for direct cortical imaging and its relevance for human cerebrovascular disease (including small vessel disease and vascular dementia) was recently highlighted.⁵² We show new insights into the kinetics of hypoperfusion, BBB dysfunction, cell damage and free radical formation after vascular occlusion, and propose that clinical assessment of BBB permeability may help to delineate tissue at risk in patients.

DISCLOSURE/CONFLICT OF INTEREST

The authors declare no conflict of interest.

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