The Temporal Dynamics of Poststroke Neuroinflammation: A Longitudinal Diffusion Tensor Imaging–Guided PET Study with $^{11}\text{C}$-PK11195 in Acute Subcortical Stroke

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Animal experiments suggest that 2 different types of activated microglia (AMG) cells occur in the brain after a stroke: local AMG in the area of the infarct and remote AMG, which occurs along affected fiber tracts. We used $^{11}\text{C}$-PK11195 PET to image AMG in vivo after stroke in humans in a prospective longitudinal study to investigate the temporal dynamics of AMG and relate local and remote AMG activity to pyramidal tract (PT) damage using diffusion tensor imaging (DTI).

**Methods:** Eighteen patients underwent DTI–MRI, $^{11}\text{C}$-PK11195 PET, and behavioral testing within 2 wk and 6 mo of acute subcortical stroke. In 12 patients, the PT was affected by the stroke (PT group), and in 6 patients it was not (non-PT group). Standardized volumes of interest (VOIs) were placed along the PT at the level of the brain stem, semioval center, and infarct. Tracer uptake ratios (ipsilateral to contralateral) were calculated for each VOI and related to tract damage (measured as fractional anisotropy ratio) and clinical outcome. Six controls underwent the same protocol but only once.

**Results:** In both patient groups, local AMG activity in the infarct was increased initially and significantly decreased over the follow-up period. In contrast, remote AMG was detected only in the PT group in the brain stem along the affected tract and persisted during follow-up. No AMG was observed retrograde to the lesion at any time. Remote AMG activity along the affected PT in the brain stem correlated with initial PT damage as measured by DTI in the same tract portion. Local AMG activity in the infarct correlated with anterograde PT damage only at follow-up. After controlling for PT damage, initial AMG activity in the brain stem showed a positive correlation with clinical outcome, whereas persisting AMG activity in the infarct tended to be negatively correlated.

**Conclusion:** DTI-guided $^{11}\text{C}$-PK11195 PET in acute subcortical stroke demonstrates differential temporal dynamics of local and remote AMG. Activity of both types related to anterograde PT damage as measured by DTI and might contribute differently to clinical outcome.

**Key Words:** stroke; inflammation; microglia; recovery; fiber tracts

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Neuroinflammation is a complex acute and chronic reaction of the central nervous system to focal brain lesions (the biologic purpose of which is still the subject of debate) (1). One cellular component of neuroinflammation comprises activation of microglia, which are the resident macrophages of the central nervous system (2).

In ischemic stroke, inflammatory changes are seen not only in the immediate surroundings of the infarct itself but also in the remote brain regions that have fiber tract connections with the affected area (3). Microglia that become active locally, in and around the infarct, have immunohistochemical properties different from microglia activated in remote brain regions (4). Locally activated microglia (AMG) consist mainly of cells expressing major histocompatibility complex class I antigens, which are thought to act primarily as scavenger cells (5). The role of remote AMG, which predominantly express major histocompatibility complex class II antigens, is less well understood. It has been hypothesized that these cells may also play a scavenger role by removing degenerating axonal debris and might thus be a histologic correlate of Wallerian (anterograde) degeneration (7).

Once activated, microglial cells express a cholesterol transporter protein (TSPO), which was formerly known as peripheral type benzodiazepine receptor, at their outer mitochondrial membrane (8,9). PK11195 is a ligand (10) that binds to TSPO and, when labeled with the positron-emitting isotope $^{11}\text{C}$, can be used to image AMG in the brain in vivo. In recent years, $^{11}\text{C}$-PK11195 has been used...
repeatedly to map AMG in various neurologic diseases. In a 
recent cross-sectional study, we demonstrated that local and 
remote AMG can be imaged in acute subcortical stroke. We 
showed that remote AMG is specific to the damaged pyr-
amidal tract (PT) anterograde to the infarct (17).

In this paper, we present longitudinal data on the 
temporal development of AMG over a 6-mo period in 
patients with a first acute subcortical stroke and relate AMG 
activity to diffusion tensor imaging (DTI) parameters of 
fiber tract integrity and clinical outcome. DTI acquires 
information about the anisotropic diffusion of water 
molecules along white matter fiber tracts and provides 
details about tissue microstructure. Fractional anisotropy 
(FA) provides an index of the diffusion characteristics of 
water molecules preferentially directed along the axis of 
major axonal pathways. Changes in FA have been shown to 
reflect damage to the PT in patients with internal capsule 
stroke (13). Reduced FA has traditionally been interpreted 
as evidence of Wallerian degeneration and axonal loss after 
stroke (14).

In our study, we specifically addressed whether, in 
human stroke patients, changes in AMG activity over time 
were similar for local and remote AMG, degree of local and 
remote activity related to the extent of fiber tract damage, 
and initial local and remote AMG activity related to 
outcome, independent of permanent tract damage.

**MATERIALS AND Methods**

**Patients**

Eighteen right-handed patients (13 women, 5 men; mean 
age ± SD, 73.06 ± 12.99 y) were recruited, assessed, and 
scanned within 3 wk of a first-ever subcortical stroke and at 
6 mo (Table 1). The control group consisted of 6 patients 
with transient ischemic attack (TIA) (2 women, 4 men) and 
no structural lesion on the MR image. The PT group con-
isted of 12 patients (8 women, 4 men) with subcortical 
ischemic infarction affecting the PT as determined on T1-
weighted MRI (7 internal capsule, 3 basal ganglia, and 2 
semioval center). Two patients were lost to follow-up. The 
non-PT group consisted of 6 patients (5 women, 1 man) 
with subcortical ischemic infarction outside the PT. All 
groups were similar with respect to age and cardiovascular 
risk factors. All participants were recruited from the speci-
ialized Stroke Unit at the Jewish General Hospital in Mon-
treal, Canada. The study protocol was approved by the 
McGill University Institutional Review Board, and written 
informed consent was obtained from all participants.

**Behavioral Testing**

All patients were assessed for clinical stroke severity and 
were physically and neurologically examined in the acute 
poststroke phase. Stroke-related neurologic disability was 
evaluated using the National Institutes of Health Stroke 
Scale, and performance in the activities of daily living was 
assessed using the Barthel Index. The degree of handicap 
was assessed with the modified Rankin scale, and upper
limb motor function was assessed with the arm subsection (14) of the Rivermead Motor Function Test (RMFT) (Table 1).

**MRI**

T1-weighted (1-mm isotropic voxels; spoiled gradient-echo sequence; repetition time, 9.7 ms; echo time, 4 ms; and α, 12°) and diffusion-weighted images (single-shot spin-echo echo-planar sequence, with twice-refocused balanced diffusion-encoding gradients (15); 2 × 64 diffusion-encoding directions, 2-mm isotropic voxel size; 63 slices; b-value [degree of diffusion weighting] = 1,000 s/mm²; echo time, 121 ms; repetition time, 11.1 s; and generalized autocalibrating partial parallel acquisition reconstruction) were acquired on a 3-T Siemens Trio Scanner.

DTI datasets were processed using in-house MINC suite tools (McConnell Brain Imaging Centre, Montreal Neurologic Institute, McGill University, http://www2.bic.mni.mcgill.ca/). Datasets were motion-corrected and realigned to the T1-weighted anatomic scan using a mutual information-based algorithm (16). Mean diffusivity, FA, eigenvectors, and eigenvalues of the diffusion tensor were then calculated at each voxel of the 4-dimensional image. Fiber tracking was performed by the FACT (fiber assignment using a continuous-tracking algorithm) (17). The tracking was stopped if the FA was less than 0.2 or the curvature from one voxel to the next was greater than 40°. Only tracts that passed through the 2 tract–delineating volumes of interest (VOIs) in each hemisphere were retained. In the non-PT and control groups, the posterior limb of the internal capsule and cerebral peduncle were used. Because the internal capsule was affected in most of the PT group patients, the infract region itself was used as a VOI in the affected hemisphere. For unaffected hemisphere regions of interest in the PT group, infract regions were mirrored from the affected side.

The fractional anisotropy ratio ($R_{FA}$) was calculated as a ratio of the affected and unaffected hemisphere FA in the PT and non-PT groups and as left–right hemisphere $R_{FA}$ in the control group. In the PT group, tracts were subdivided into anterograde (i.e., below the stroke) and retrograde (i.e., above the stroke) portions relative to the lesion site.

**PET**

($R$)-$^{11}$C-PK11195 was synthesized from its precursor ($R$)-$^{11}$C-N-desmethyl-PK11195 (ABX) according to the method of Camsonne et al. (18), yielding 2.59–2.96 GBq (70–80 mCi) (18%–20% radiochemical yield) of the final sterile product. The specific activity was in the range of 18.5–22.2 TBq/mmol (500–600 Ci/mmol). PET scans were obtained on a Siemens ECAT HR + scanner in 3-dimensional (3D) mode (resolution at 1 cm from center, 4.4-mm radial, 5.1-mm axial). Patients were injected with 550 MBq of ($R$)-$^{11}$C-PK11195 into an antecubital vein after a 10-min transmission scan. Dynamic emission data were acquired over 60 min (19). A filtered backprojection algorithm, supplied by the manufacturer, with a Hanning filter cutoff frequency of 0.4 cycles per pixel was used for reconstruction. After correction for scatter, attenuation, and random coincidences, an image volume comprising 21 frames with matrix dimensions of 128 × 128 × 63 voxels and a voxel size of 2.06 × 2.06 × 2.43 mm was obtained.

PET data analysis was performed in VINCI (version 2.55; Max Planck Institute for Neurological Research). Individual PET data were coregistered to the T1-weighted MR images (20), and fiber tract images were overlaid onto the anatomic MR images. Because tractography identifies only the remaining intact fibers on the affected side, tracts were significantly smaller on the affected side in the PT group. Thus, we did not use the tracts themselves for the analysis but placed a standard set of elliptic 3D VOIs along both PTs, manually adjusted these VOIs to the individual size of the brain stem (comprising 1 VOI for pons and 1 for mid brain) and semioval center on the unaffected side, and copied the VOIs to the affected tract. This procedure ensured that nearly identical VOIs were used on both sides, despite smaller residual tracts on the affected side. These standard VOIs would extend about 2 mm beyond the outlined tract to cover the entire cerebral peduncle and thus also include tracer uptake in the immediate surroundings of the tract (Fig. 1). One further ellipsoid region was placed in the infract and mirrored about the interhemispheric fissure to obtain the counterpart region in the unaffected hemisphere. In patients with TIA the infract regions were placed in the basal ganglia at the level of the internal capsule.

Uptake ratios (URs) between corresponding VOIs (affected side divided by unaffected side) were calculated from an average image of the last 40 min. Ratios greater than 1 indicate higher uptake on the affected side. These URs were compared within and between groups using 2-way ANOVA with the Student–Newman–Keuls (SNK) statistic for posthoc analysis at a corrected type I error level of 0.05. The use of 1 average image of the last 40 min for analysis was possible because the URs of averaged time–activity curves (time–activity curve for the affected side divided by time–activity curve for the unaffected side) were stable for a low- (semioval center) and high-affinity (infract) VOI at 20 min after injection.

**FIGURE 1.** Placement of standardized 3D ellipsoid VOIs along tracings of PT derived from diffusion tensor images. VOI for brain stem (BS) consists of 2 ellipsoids for mid brain and pons. One 3D ellipsoid was used for semioval center (OC).
Within- and between-group differences were tested using repeated-measures ANOVA as appropriate, corrected for multiple comparisons (SNK statistic, \( P = 0.01 \) corrected). Linear regression analysis was performed between \( R_{FA} \) values and URs in the brain stem and infarct. Initial URs correlated with clinical outcome using Pearson product moment correlation.

**RESULTS**

**Regional \(^{11}\text{C-PK11195 Uptake}\)**

The tracer UR in the infarct (UR\(_{inf}\)) was significantly increased at the initial measurement in both the PT and the non-PT groups, as compared with the control group (\( P < 0.01 \)), with no difference between these 2 patient groups. During the follow-up period, tracer uptake decreased significantly in both patient groups and was no longer different from that of the control group (2-way repeated-measures ANOVA, SNK test for multiple comparisons; Fig. 2). Tracer uptake along the PT was increased in the PT group but not in the non-PT and control groups and only distal to the lesion (UR at the level of the brain stem [UR\(_{brain\ stem}\]), indicating that remote AMG activity is specific for the affected PT below the infarct (Fig. 3A; 2-way ANOVA \( P < 0.01 \) corrected)). This considerable anterograde increase persisted over the 6-mo period (Fig. 3B; ANOVA, \( P < 0.01 \) corrected) and was not significantly different from the initial measurement (2-way repeated-measures ANOVA, \( P > 0.05 \)). No changes were observed in retrograde tract portions (semioval center).

Differences in URs were also reflected in absolute activity concentration differences in Bq/cm\(^3\) between VOIs on the affected side and VOIs on the unaffected side (data not shown).

**PT Damage and Correlation with \(^{11}\text{C-PK11195 Uptake}\)**

The \( R_{FA} \) was lower for the entire tract (total \( R_{FA} \)) and for the tract portions above (retrograde) and below \( (R_{FA\text{-below}}, \text{anterograde}) \) the infarct in the PT group, as compared with the non-PT group and controls at both times, thus indicating permanent tract damage in the PT group (2-way repeated-measures ANOVA, SNK statistic, \( P < 0.01 \)). No significant change in \( R_{FA} \) was observed over time (Table 1).

At the initial measurement, the UR\(_{brain\ stem}\) for \(^{11}\text{C-PK11195}\) was significantly negatively correlated with \( R_{FA\text{-below}} \) (\( r = -0.70, P = 0.019; \) Fig. 4), indicating higher microglial activation with increasing tract damage (decreasing \( R_{FA\text{-below}} \)). A similar relationship was found for the follow-up measurement (\( r = -0.72, P = 0.012; \) Fig. 4).
There was no significant relationship between tracer URinf and tract damage (either entire tract or any of its portions) for the initial measurements. At follow-up, however, a highly significant correlation between persisting URinf and RFA-below was found ($r = 0.92$, $P < 0.01$; Figs. 4 and 5).

**Correlation of Initial Parameters with Clinical Outcome**

Motor function as measured by the RMFT was initially more impaired in patients in the PT group (rank sum test, $P = 0.041$) than in the non-PT group. PT group patients had significantly higher RMFT scores at follow-up than initially (Wilcoxon test, $P = 0.047$); no significant change was observed in the non-PT group. In the PT group, initial tract total $R_{FA}$ correlated significantly with initial RMFT ($r = 0.92$, $P < 0.001$), and follow-up total $R_{FA}$ correlated significantly with RMFT scores at follow-up ($r = 0.78$, $P = 0.012$).

To further investigate the impact of initial local and remote AMG on outcome, we correlated initial URinf and URbrain stem with RMFT at follow-up. No correlation was found between initial URbrain stem and motor function. When controlling for permanent tract damage (total $R_{FA}$ at follow-up) using partial correlation, the initial URbrain stem showed a significant positive correlation with follow-up RMFT score ($r = 0.67$, $P = 0.035$). The initial URinf had a trend toward a negative association with motor outcome ($r = -0.63$, $P = 0.053$). This trend persisted when controlling for permanent tract damage (total $R_{FA}$ at follow-up) using partial correlation ($r = -0.62$, $P = 0.052$).

**DISCUSSION**

This is the first—to our knowledge—prospective, controlled longitudinal study describing the temporal dynamics of AMG and its relationship to morphologic PT damage in humans with subcortical stroke using $^{11}$C-PK11195 PET and DTI. Although microglial activity in the infarct decreases during recovery, remote activity in the brain stem along the affected tract persists. Microglial activation in general seems to be related only to anterograde PT damage but again with different temporal dynamics: in the brain...
FIGURE 5. (A) Microglial activation in patient with small subcortical infarct and good recovery. Initial activated microglia in infarct decreased over 6 mo (white arrows), whereas microglial activation in brain stem persisted (red arrows). DTI showed decreased FA primarily in infarct (blue arrows) and less along tract at level of cerebral peduncles (yellow arrows). (B) Patient with complete transection of PT and poor recovery. Microglial activity in infarct decreased but still persisted after 6 mo (white arrows), as did activity in brain stem (red arrows). FA decreased in area of infarct (blue arrows) and along tract in cerebral peduncle (yellow arrows). Microglial activity in patient in whom PT was not affected (C) decreased over 6 mo (white arrows). No tracer uptake at level of brain stem was observed, and FA along tracts was not decreased.
stem, remote AMG related to PT damage during the entire observation period, whereas microglial activity in the infarct related to PT damage only after 6 mo.

**Does 11C-PK11195 Measure Microglial Activity in Stroke?**

11C-PK11195 has been widely used for imaging of AMG during the past 20 y (21). Early studies in animal models of ischemic stroke, using autoradiography in combination with immunostaining for macrophage cell lines, demonstrated that the 11C-PK11195 signal associated with ischemic lesions originated primarily from macrophages (22). A more recent immunohistochemical study using antibodies that directly bind to the TSPO receptor in postmortem human brains confirmed that the expression of TSPO in the normal brain is minimal and that microglia and macrophages are the primary cell types exhibiting an increased number of TSPO binding sites in human pathologies (8). Astrocytes can also express TSPO in degenerative disease models (27) or in the infarct, where they contribute to scar formation. A recent small-animal PET study in rats using a new TSPO ligand directly compared immunohistology with imaging data and found that microglial and not astrocyte migration best correlated with the time course of tracer uptake. The authors concluded that either expression of TSPO is higher in microglia than in astrocytes, even at late times after ischemia, or different glial cell types express different TSPO subtypes (24). The contribution of astrocytes to the PET signal thus awaits further clarification.

Although there are some limitations to the use of 11C-PK11195 as a radiotracer for microglia imaging (signal-to-noise ratio, unspecific binding), the most promising alternatives (25) have so far been tested only in a few healthy subjects (26,27), so the use of these alternative radiotracers in ischemia or other brain pathologies has not yet been established. In this study, we addressed the limitations of 11C-PK11195 in 2 ways. First, given the specific hypotheses (increase of tracer uptake along the PT in the brain stem and semi-oval center), we have chosen a VOI approach, which is more robust to fluctuations in signal-to-noise ratio. Second, we used URs relative to mirrored regions in the unaffected hemisphere to avoid the use of a single reference region, thus accounting for a possible varying tracer binding in the unaffected hemisphere. Even if there were some unspecific binding in those mirrored regions, it would only bias our results in a conservative direction.

**AMG in Infarct**

The time course and cellular physiology of AMG in ischemia has been extensively studied in animal models of stroke. About 3–5 d after 1 h of transient focal ischemia, microglial activity has been found to peak in and around the area of the infarct in a mouse model, persisting at relatively high levels (28). In a recent imaging study using a new TSPO receptor ligand with a 2-h transient focal ischemia model in rats, peak activities were observed after 9–11 d (24). Local AMG can produce toxic substances that may ultimately promote delayed death of neurons around the infarct (29). However, AMG around the infarct may also be indirectly neuroprotective by neutralizing the effect of neutrophils (30) and removing debris from the infarct site (5). More recently, it has been demonstrated that AMG can also produce BDNF (brain-derived neurotrophic factor) (31) and thus actively contribute to neuronal regeneration.

Previous case series with 11C-PK11195 PET in human stroke showed results comparable to those of animal studies with respect to the early occurrence of the microglial response (~day 3), which persisted in some subjects up to 150 d (32). The specificity of those findings, however, remained unclear because of the lack of controls. In our study, an early increase (3 d in the earliest of our patients) was observed in the infarct region of all stroke patients. The initial presence of AMG appeared to be a more-or-less stereotyped reaction and was not significantly different between the PT group and non-PT groups. This observation is in line with a recent 11C-PK11195 study in a rat model of permanent ischemia, in which the extent of microglial activation also was independent of infarct location or size (33).

Over the 6-mo follow-up period, the UR decreased in all stroke patients. Although on average this activity was no longer significantly different from that in the controls, some variability of 11C-PK11195 URs in the infarct remained (Fig. 2). Data from animal experiments on such extended periods are limited. In a recent study in rats, Keiner et al. also described a decrease of microglial activity in the perilesional zone at 42 d after permanent photothermibotic ischemia (34).

**Remote AMG**

In contrast to the infarct region, 11C-PK11195 URs in the brain stem were significantly increased initially and at follow-up but only in those patients in whom the PT was affected by the stroke (Fig. 3). These results expand on previous observations from uncontrolled case series (35), because they demonstrate that increased URs in the brain stem are indeed specific for the damaged tract and not an unspecific reaction to the ischemic event per se. Our results suggest that this increased remote uptake has temporal dynamics different from AMG at the infarct site. Remote microglial activity has previously been described in animal models to occur around day 2 after the infarct and maximize around 7 d after ischemia (6). However, in single case studies of human stroke, ipsilesional thalamic 11C-PK11195 binding was detected several months after the event (35). In our study, the URs at follow-up were not significantly different from initial URs, thus indicating that microglial activity along the damaged tract is relatively constant over the 6-mo follow-up period. Some tracer uptake was also observed in the unaffected hemisphere (e.g., the thalamus in Fig. 5). This uptake may correspond to an increased level of inflammation related to microangio-pathy secondary to diabetes, hypertension, or age (36).

Larger studies with stratification of risk factors and a tracer
with less unspecific binding than PK11195 are needed to verify this observation. These results again support the choice of TIA patients as controls; these TIA patients were comparable to the patient group with respect to these factors.

Relationship to Fiber Tract Damage

FA measurements of fiber tracts are generally regarded as an indicator of tract damage (37). We have shown that RFA within the first weeks correlate with clinical stroke severity and outcome after 6 mo. This finding seems to indicate that initial RFA are a good indicator of the extent of acute and long-term ischemic damage. A decrease in FA distal to the lesion after the first few weeks has been described by some authors (12) and interpreted as a correlate of Wallerian degeneration (38). We found a significantly lower RFA initially and at follow-up for the entire tract and for the tract portions above and below the infarct. 11C-PK11195 uptake, whether locally in the infarct or remotely in the brain stem, related to fiber tract damage in only the portion below the infarct. Again, remote and local AMG show a different association to tract damage: although remote AMG related to RFA_below initially and at follow-up, AMG in the infarct was highly correlated with RFA_below only after 6 mo (Fig. 4). This result suggests that the extent of acute (mostly ischemic) damage to the tract in the first days after the stroke determines the intensity of remote, anterograde microglial activation and may thus be related to the amount of axonal damage that will eventually lead to Wallerian degeneration. This interpretation is in line with data from an animal study in focal intracerebral hemorrhage (39).

In contrast, AMG in the infarct at the initial measurement is not related to anterograde tract damage per se. Only over time, when degeneration sets in (which is reflected by the RFA_below at follow-up), does persisting AMG significantly correlate with anterograde tract damage. Although AMG in the infarct decreases over time on average (Fig. 2), there are still patients with increased URs 6 mo after the ischemia. In those patients, persistent activity seems to have caused more anterograde degeneration.

The 2 patients in Figure 5 illustrate persistent activity. A patient with partial damage to the PT (Fig. 5A) exhibits microglial activation in the infarct, which vanishes over time. Damage to the tract is mainly limited to the area of the infarct in the corresponding FA image; the tract itself is less affected. In contrast, AMG persists in a patient with complete transection of the PT at the level of the internal capsule (Fig. 5B) and poor outcome. Here the FA image already shows tract damage at the level of the brain stem. In both patients, remote AMG is found at both times. No AMG is observed in the brain stem of a patient in whom the PT is not affected (Fig. 5C).

AMG and Clinical Outcome

The question of whether increased AMG activity is beneficial or detrimental to poststroke recovery remains unanswered (40). A recent stroke-recovery study in rodents observed better outcome in animals with lower AMG activity in the perinfarct zone (34). To explore these relationships further, we asked the question of whether measurement of local or remote AMG early after stroke might provide any additional information on stroke outcome beyond what already obtained through FA measurement. We thus performed a partial correlation analysis between initial URs and clinical outcome after 6 mo while controlling for damage to the entire tract as measured with RFA. Interestingly, there was a trend toward a negative partial correlation between initial microglial activity in the infarct and outcome as assessed with the RMFT, whereas remote AMG was positively correlated with outcome. The results of this exploratory analysis may suggest that beyond ischemic tract damage initial microglial activation in the area of the infarct might influence outcome negatively, and initial remote AMG might be regarded as a positive influence. However, the relative impact of each of these processes and their exact role for stroke recovery remain to be determined.

CONCLUSION

The results of this study provide the first, to our knowledge, in vivo evidence for a relationship between microglial activity and fiber tract integrity in human subcortical stroke. We demonstrated that perinfarct and remote microglial activity not only follow different time courses during a 6-mo period after subcortical stroke but also are differentially related to anterograde fiber tract damage and might contribute differently to clinical outcome. In vivo microglia imaging combining PET and DTI may thus prove useful in the assessment of complex interrelationships between neuronal damage and poststroke neuroinflammation and provide surrogate markers for testing therapeutic strategies to modulate microglial activity toward favorable recovery at the transition from animal experiments to clinical trials.

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REFERENCES


NEUROINFLAMMATION IN SUBCORTICAL STROKE • Thiel et al. 1411


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