How can sodium MRI techniques help us understand acute stroke?

This article addresses the potential usefulness of sodium MRI in the acute phase of stroke, asking whether the additional time required to acquire high-quality sodium images is justified in the time-critical minutes following the presentation of a patient with stroke symptoms. It begins with a description of the pathophysiology of stroke and the implications of the increasing bioenergetic failure on the sodium content in tissue. Recent studies that have aimed at imaging the subtle changes in this sodium content in stroke patients are then reviewed, followed by experiments in animal models of stroke, which circumvent some of the limitations of the human studies. Finally, steps that will probably be required to translate these latest model findings into human studies are discussed, including new MRI techniques that may provide a boost in signal and allow for the introduction of relaxation-time contrast and quantification of the sodium concentration.

KEYWORDS: penumbra  sodium MRI  sodium perfusion mismatch  sodium quantification  stroke  tissue sodium

The aim of this article is to provide a critical overview of the current use of sodium (Na)-MRI in the acute phase of stroke, focusing initially on the relatively limited studies conducted to date in humans and then moving on to describe the more numerous studies that have been carried out in animal models of stroke, which despite the inherent limitation of such models, particularly when translating findings to the human system, nevertheless may indicate areas for future focus in human studies. It can be argued that, despite the drastically more complicated cerebral vascular system in the human compared with that in the rat brain, for example, at a cellular level both human and rat brain systems are not so dissimilar and cellular bioenergetic failure and the consequent change in Na signal detectable via MRI can thus be expected to manifest in both systems in a similar manner. The main difference is in the time at which the bioenergetic failure occurs in brain regions downstream from a blockage: in rat models, most of the damage occurs relatively quickly, with the stroke lesion reaching its maximum extent within 4–6 h after the stroke was induced. In the human brain, by contrast, the vastly more sophisticated vascular system, which incorporates a higher degree of redundant blood supply to most areas (the so-called collateral supply), the situation is more complicated with a slower evolution of the stroke lesion extending out to 48 h or more, suggesting a lengthened time window in which to intervene therapeutically. Thus, changes in the Na MR signal observed in animal models, whether due to changes in Na concentration or perhaps relaxation-time-related signal changes specifically introduced into the images, should be translatable into human studies. This article sets out the current state of the art in this area, and highlights the challenges that will need to be overcome to allow for routine clinical imaging of acute stroke patients.

Background
After attempts to use Na-MRI in the 1980s met with limited success, the field is currently enjoying a renaissance driven primarily by technological developments in MR technology that are making it possible to explore the many possible clinical applications of Na-MRI, some of which have been postulated since the early days of MRI but have heretofore remained tantalizingly beyond reach. Indeed, Na spin-density-weighted images with reasonable signal to noise ratio (SNR) and spatial resolution can now be obtained in the brain in less than 10 min using standard 3 T clinical imaging platforms [1], and a growing number of groups worldwide are working to push the boundaries of this exciting technique ever further. Examples of clinical application areas that have been investigated to date include cerebral stroke [2–4], brain tumor [5,6], breast tumor [7], cardiac infarction [8], Alzheimer’s disease [9], multiple sclerosis [10], kidney [11,12] and osteoarthritis in cartilage [13,14], while the first whole-body Na images were recently presented [15].
In addition, considerable effort is going into the extension of the Na-MRI technique beyond mere spin-density weighting, to explore the introduction of relaxation-time contrast and indeed relaxometry mapping and the use of multiple quantum filter techniques to separate intra- from extra-cellular Na compartments in tissue, while future developments may well involve Na diffusion-weighted imaging and beyond. The resurgence of Na-MRI has come about primarily as a result of the maturation of MR technology, encompassing the availability of high-field strength systems (≥ 3 T), improvements in system electronics and radiofrequency (RF) coil design concepts, and advanced imaging pulse sequences, which combine rapid acquisitions with high sensitivity to the short T₁-relaxation time components of Na nuclei encountered in vivo.

The change in Na concentration in infarcted tissue, that is, tissue that has died following bioenergetic failure due to the interruption of blood supply to the area, typically following a stroke, has been known for many years, and indeed early attempts to study both cerebral and cardiac infarction in humans via Na-MRI were largely proof-of-concept and revealed little of use for acute patient imaging. In more recent years, a growing number of studies in preclinical animal models of cerebral stroke have been reported in the literature, to the level where a picture of the potential usefulness of Na-MRI for acute stroke imaging is now emerging. The critical role of Na⁺ in maintaining healthy cellular function, followed by the physiological link between Na concentration and infarction, is outlined in the next section.

**Link between Na & stroke**

The potential of Na-MRI for many disease investigations is underpinned by the critical role of Na in many cellular functions such as the regulation of mitosis, cellular proliferation, the mediation of Na/calcium exchange, and energy metabolism via the Na⁺/K⁺ ATPase pump function located in the cell membrane. Indeed, the large Na⁺ concentration gradient across the cell membrane established by the Na⁺/K⁺ ATPase, which maintains the intracellular concentration at approximately 10 mmol/l and the extracellular Na concentration at approximately 140 mmol/l, is essential for the generation and propagation of action potentials, cell volume regulation and other cellular homeostatic and regulatory functions. Disease processes that alter this energy-dependent pump function will thus lead to an altered tissue Na content, which may be detected via Na-MRI techniques.

The mortality and morbidity associated with stroke are well documented: it is the third leading cause of death and the leading cause of disability in developed countries [16]. Despite much research in the area of neuroprotectants and therapies, only one approved treatment currently exists, intravenous thrombolysis with recombinant tissue plasminogen activator (rtPA). However, it is only approved for administration within 4.5 h of symptom onset, a time window based on epidemiological studies of large cohorts of patients, which effectively excludes its use in many patients for whom the time from onset is unknown or exceeds the time window (although intrarterial thrombolysis out to 6 h is currently performed in many institutions). Its use is further restricted owing to other factors such as the risk of intracranial hemorrhage associated with rtPA (7%) [16], such that the estimated uptake of rtPA is less than 10% in stroke patients [17]. It is likely that a more tailored approach to the use of this treatment in individual patients would be appropriate, for example allowing for an extension of the time window during which it could be administered, and indeed recent studies have suggested that there may be benefit to using rtPA out to 6 h [18]. However, such a tailored approach requires a reliable method for assessing the potential for benefit of rtPA to an individual, and it is here that Na-MRI, coupled with other MRI techniques, may have a role to play.

Ischemic stroke is a catastrophic event in which the blood supply to a part of the brain is restricted, and oxygen and glucose supply to cells within the affected region is severely limited. The loss of nutrient supply to cells in turn compromises energy metabolism within the stroke-affected area, leading to a rapid loss in ATP and hence in ionic homeostasis. One immediate consequence of this is a reduced efficiency of the Na⁺/K⁺ pump function and a resulting increase in the intracellular Na concentration. This in turn leads to edema and eventually to cell membrane rupture. To stratify patients who might benefit from rtPA treatment, conventional MRI techniques employed in the acute phase of stroke have attempted to identify the presence of so-called ‘penumbral’ tissue regions, which are hypoperfused yet still viable using the so-called ‘perfusion–diffusion mismatch’, wherein the mismatch between the ischaemically injured tissue and hypoperfused tissue is determined from diffusion-weighted imaging (DWI) and perfusion-weighted ¹H-MRI imaging (PWI),...
How can sodium MRI techniques help us understand acute stroke?

Tissue in such penumbra regions are at risk of infarction, but can be salvaged by a timely reperfusion of the affected tissue, and hence the presence of such at-risk tissue in a stroke patient would give confidence for a potential benefit for the administration of rtPA.

The DWI technique is sensitive to the molecular diffusion of water molecules in tissue, and indeed can be used to obtain a quantitative estimate of the diffusion coefficient (the apparent diffusion coefficient [ADC]). The cytotoxic edema, which occurs within minutes of the ischemic onset results in the restriction of the diffusion of water molecules within the affected tissue, which can be easily detected in an ADC map. The PWI technique tracks a bolus of paramagnetic contrast agent as it perfuses through tissue and provides a semi-quantitative measure of the regional cerebral blood flow, and thus may be used to determine the degree of underperfusion caused by the blockage. However, despite early optimism relating to the perfusion-diffusion mismatch and studies that have shown that thrombolytic treatment outcomes are improved in patients selected based on an MRI diagnosis [19], it is now clear that this approach leads to a lack of precision in the separation of still-viable from irreversibly damaged ischemic core tissue. For example, large deviations have been observed between areas with decreased ADC compared with infarct determined by the gold standard technique of histology after stroke [20], while ADC values have also been observed to normalize in areas of permanently damaged tissue during the subacute phase [21], suggesting that the lesion delineated by DWI does not provide a definitive indicator of infarcted tissue in the acute stroke phase. Furthermore, definitive thresholds of blood flow for penumbra or infarct have not been identified, and hence the area of hypoperfusion revealed by PWI may overestimate the volume of tissue that is realistically ‘at-risk’, owing to the inclusion of areas of benign oligemia. Overcoming this problem will require the establishment of such thresholds, although it is known that they depend on the duration of ischemia and any reperfusion that has occurred, whether thrombolysis-induced or spontaneous [22]. New approaches to the analysis of perfusion-weighted data may moderate the severity of the overestimation [23].

The Na-MRI technique provides a measure of the tissue sodium concentration (TSC), which represents an average of the Na concentration in the intra- and extra-cellular tissue compartments weighted by their respective volume fractions. For example, in brain tissue the TSC is typically 45 mmol/l, considering an intracellular volume fraction of approximately 70% and an extracellular volume fraction of 30% (the latter comprising a 5% vascular space and a 25% extracellular extravascular space) Following an ischemic insult to the tissue and the failure of the Na+/K+ pump, the intracellular Na concentration gradually increases above its nominal 10–15 mmol/l level, while the extracellular concentration, buffered by the body, is maintained at 140 mmol/l. This results in a net increase in the TSC and has been measured in both animal model studies [24–28] and human stroke patients [2–4,29].

Challenges with imaging of Na nuclei

The difficulty with Na-MRI lies primarily with the lower concentration and MR sensitivity of Na nuclei compared with hydrogen nuclei in vivo, resulting in signal levels approximately 20,000-times lower than that obtained using conventional 1H-MRI techniques. In addition, the resonant frequency of Na nuclei is approximately one-quarter that of hydrogen nuclei, and hence broadband MRI systems with dedicated RF transmit amplifiers and RF coils for transmission and detection of the Na signal are also required; although available for clinical 3 T systems, at present these hardware options are not typically installed as standard. Indeed, the development of new RF coil designs has contributed to the increased SNR achievable in Na-MRI experiments and remains an active area of research. For human brain imaging, transceiver volume resonators based on a birdcage design are typically used; examples include single-tuned resonators for 23Na frequencies only, necessitating a swap of the RF coils to acquire 1H images [3,6], or dual-tuned to allow for 1H and 23Na imaging without disturbing the patient in the scanner and losing valuable time [30,31]. However, such dual-coil configurations invariably compromise the SNR on one channel, usually selected to be the 1H channel, and the effect this may have on the quality of the images typically acquired as part of a clinical stroke protocol, for example involving time-of-flight angiography or echo planar imaging-based diffusion and perfusion measurements, have not been reported. Transceiver surface coils have also been used for imaging outside the brain, for example in heart [32] and breast imaging [33], although quantification is extremely difficult due to inhomogeneous transmit RF fields from such coils. A more recent development for Na-MRI has
seen the separation of the transmit and receive functions, with a volume resonator used for the former and a surface coil for the latter function [35]. This approach is particularly suited to applications outside the brain where detectors can be designed to optimally fit the anatomy of interest (e.g., breast, kidney), thereby optimizing the SNR, while the volume transmitter provides a uniform flip angle excitation that facilitates accurate quantification of the TSC. The development of phased array coils for the brain should extend these advantages to stroke imaging. For rodent brain imaging, such a dual-coil approach has been found to be essential for accurate TSC quantification [26].

The lower SNR for Na in vivo can be partially compensated by the use of imaging sequences with short repetition times (TR), since the longitudinal T1 relaxation time for Na nuclei (measured to be 36 ms in the human brain at 4.7 T [34]) are much shorter than for hydrogen. However, a further complication for Na-MRI arises from the quadrupolar nature of the Na nucleus due to its spin 3/2 property, which owing to its interaction with local electric field gradients from neighboring nuclei in the tissue leads to a biexponential transversal T2 relaxation, with fast and slow components (T2 ~1–2 ms and 15–20 ms in the brain, respectively) with relative proportions of 60% and 40%, respectively. To acquire signal from the fast component, one must use a short echo time imaging sequence; examples of such sequences that have been used for Na imaging in stroke include 3D projection-reconstruction (3DPR) [35], twisted projection imaging [36] and density-adapted 3DPR [31].

Clinical stroke Na-MRI studies

Acute stroke patients are often quite sick and there are significant logistical difficulties involved in carrying out Na imaging experiments in this patient cohort, not least the maxim that ‘time is neurons’, which makes it ethically difficult to justify the experimental imaging of patients which may postpone any available therapy options. As a result, only a handful of such studies appear in the scientific literature to date, and many patients recruited to these studies underwent Na imaging after a thrombolytic treatment was administered, so any reperfusion effects caused by these treatments (or indeed due to spontaneous reperfusion) represent a potential confound to the results obtained. Furthermore, patient numbers on these studies have been quite low, which, coupled with the inherent heterogeneity of human strokes, renders it difficult to draw definitive conclusions from existing studies.

The first study describing Na imaging in the brain was in 1988 and although increased signal was seen in patients with herpes simplex encephalitis (due to the resulting edema) and tumors, no signal change was detected in the single stroke patient imaged owing most likely to the very poor image quality [37]. The first dedicated study of Na-MRI in stroke was reported by Shimizu et al. in 1993, who imaged seven patients with confirmed stroke (three within 14 h of symptom onset and the rest at 47–82 h after onset) using a 1.5 T scanner [38]. The authors of this pioneering study reported no change in the Na signal in the acute phase (that is, out to 13 h; n = 3), however, a ‘marked signal increase’ (not specified by the authors) was detected in the ischemic area at 38 to 82 h (n = 4). It is likely that the poor quality images achieved in this early study, owing to the use of a spin echo sequence with a long echo time of 13 ms, producing significant T2 signal decay, resulted in the inability to detect any change in Na signal in the acute phase. In a study by Thulborn et al. in 1999, Na images were acquired on 42 patients with confirmed stroke, 11 of whom were imaged within 24 h of symptom onset [29]. Na imaging was performed on either a 1.5 T or 3 T scanner, using a dual-tuned 23Na/H volume resonator, with all acute patients imaged using the 1.5 T scanner. The quantified TSC was found to be elevated in the infarcted region by at least 25% in all patients (where the infarcted regions were defined by analysis of corresponding DWI and PWI data), although no clear pattern of TSC increase could be discerned in the hours and indeed days after stroke onset. The existence of a threshold TSC value, above which irreversible tissue damage would inevitably occur even following a therapeutic intervention, was suggested by the authors who nonetheless stated that more extensive experimentation using animal models would be required to define such a threshold.

A study in 2009 succeeded in recruiting 21 stroke patients with imaging times as early as 4 h and out to 104 h after the known stroke onset time, with repeat scans performed on ten of these patients [4]. All 1H-MRI was performed on a 1.5 T scanner, with the patients transferred to a 4.7-T system for the Na-MRI imaging, where a single-tuned resonator was used together with an imaging sequence termed ‘sodium projection acquisition in the steady state’ to give high SNR images with 10-min image acquisition times. Clear Na signal increases, relative
to the contralateral side, were observed in all patients in regions exhibiting reduced water mobility (as determined from ADC maps), with a modest increase of < 10% in patients imaged within 7 h of stroke onset (n = 5), followed by a more rapid increase out to 18–20 h, eventually reaching a plateau of 69 ± 18% after approximately 72 h. The regions of interest used in this analysis was presumed to represent infarcted tissue, and further increases in the relative Na signal of 76 ± 20% were observed in follow-up scans from 88–227 days. The ADC maps and Na images from one patient on this study who was imaged at three time-points are presented in Figure 1, demonstrating the increase in Na signal (45% increase over contralateral side) between the images acquired at 4 and 26 h after stroke onset, with a further increase (75% relative increase) at 91 days.

The most recent human stroke study reported in the literature was carried out by Tsang et al., who aimed to investigate Na signal changes in the PWI–DWI mismatch area during the acute stroke phase [2]. Nine patients were imaged 4–32 h after symptom onset, with follow-up scans performed on four, again using a 4.7 T scanner with a single-tuned volume resonator for the Na imaging. In patients imaged 4–7 h after onset, no increase (relative to contralateral homologous tissue regions) in the Na signal was observed in core tissue (where the region of core tissue was defined as the DWI-delineated lesion), although by 17–32 h a significant increase in this region was measured, with an average increase of 45 ± 26% across the nine patients. However, no correlation was found between this increase and the corresponding degree of hypoperfusion in the core regions. Furthermore, no change in Na signal was detected in regions defined by the DWI–PWI mismatch in any of the patients investigated. Data from one patient in this study is presented in Figure 2, where a Na signal increase is evident at 25.5 h but not at 4 h post-symptom onset.

Preclinical stroke Na-MRI studies

The few human Na-MRI stroke studies conducted thus far suffer from a lack of temporal data on the stroke evolution, with at most two follow-up scans performed on a subgroup of patients at best, while the heterogeneity of stroke progression evident between patients makes it difficult to draw firm conclusions as to the utility of Na-MRI in the acute phase. The use of preclinical models of stroke have a number of

![Figure 1. Representative slices from a patient imaged at three time points after stroke onset.](image)

The images show no increase in Na signal (relative to the contralateral side) at 4 h in a region of interest drawn to correspond with the ADC-defined lesion, but with increases of 45 and 75% at the later times of 24 h and 91 days, respectively. ADC: Apparent diffusion coefficient. Reproduced with permission from [4].
advantages in this regard: the ability to control the severity and extent of the ischemic damage; precise knowledge of the stroke onset time; the ability to acquire temporal data in order to follow the evolution of the stroke throughout the acute phase; and the possibility to histologically verify ischemic tissue damage. The earliest experiments in this field date from 1983 in the cat [39] and 1985 in the rat [40], where increases in the Na signal were detected 9 and 3 h after the strokes were induced, respectively. However, the relatively poor image quality of these studies deterred follow-up studies for many years, with a resurgence in activity in the last decade with the appearance of papers describing studies in a variety of animals, primarily rats, rabbits and primates. In these studies, a variety of stroke models have been used, including: the intraluminal thread model [27,41], direct middle cerebral artery occlusion with combined bilateral common carotid artery occlusion in the rat [24], and autologous embolic middle cerebral artery occlusion models in the rabbit [28] and nonhuman primate [3,29,42]. These models induced different severities of ischemic insult, depending on the addition of common carotid artery occlusion, the level of collateral blood supply in the model and the neuroanatomical location of the region of interest (e.g., ischemic core vs penumbra).

Nevertheless, a common finding of these studies was a gradual increase in the Na signal within core tissue, with rates of increase depending on the species: approximately 22–25 %/h in rats, approximately 12%/h in rabbits and ~5–13%/h in monkeys (compared with ~5%/h in humans [2,4,29], although it should be noted that an explicit linear increase in humans has not been verified thus far). This pattern of smaller rates of increase in the larger mammals most likely reflects the increasing degree of collateral supply in their brains, a feature that has complicated the translation of findings to human studies and perhaps contributed to the failure in clinical trials of several therapies and neuroprotectants, which showed promise in rodent studies.

The apparent linear increase in Na signal measured in core tissue has led to the idea that Na measurements in the acute stroke period may be used to determine the stroke onset time by extrapolating back in time, with a reported accuracy of ± 4 min in a rat model [24]. It was further hypothesized that there may exist a threshold TSC value in the vicinity of 70 mmol/l above which tissue is irreversibly infarcted, but below which tissue could recover if reperfused sufficiently, accompanied by a return of the TSC value to normal levels [29]. However, this hypothesis has not been verified to date; in a recent perfusion study in a monkey model of ischemia, the initial increase in the Na signal in the core tissue was stopped following a reperfusion of the affected brain regions, although no subsequent decrease was measured [42]. However, only three successful reperusions were achieved, while the limited spatial and temporal data acquired may have obscured any further changes.

In a recent study from my group, high temporal and spatial resolution quantified TSC data was obtained in a rat stroke model which showed regionally disparate behaviors in the TSC increases: while the TSC increased immediately in core tissue, delays to TSC increase of up to 4 h were measured in presumed penumbra tissue regions (Figure 3) [27]. In all cases, once the TSC was observed to increase, the rate of increase was the same across all areas irrespective of the delay time to increase, while areas exhibiting any increase in TSC above normal were correlated with histologically verified infarction, suggesting that infarction of tissue occurs once the TSC increases above normal levels. Of further note, in this study there was a decrease of approximately 4 mmol/l measured in presumed penumbra tissue; a follow-up study where Na, diffusion- and perfusion-weighted data were correlated with

Figure 2. Perfusion, diffusion and sodium data from a representative slice of a stroke patient imaged at two time-points post-symptom onset. The DWI hyperintense region is indicated by the dotted outline, with the PWI–DWI mismatch tissue by the solid outline. Note the lack of sodium signal change at 4 h, which increased to 1.46 (relative to the contralateral side) by 25.5 h in the core region only.

DWI: Diffusion-weighted imaging; PWI: Perfusion-weighted imaging. Reproduced with permission from [2].
histopathology, which allowed for accurate penumbra identification, detected for the first time a decrease to $90 \pm 8\%$ of the contralateral Na signal ($n = 6$) in penumbra tissue, with the maximum decrease occurring $106 \pm 60$ min after stroke onset [43]. Representative images from one rat showing the decrease in the Na signal in the penumbra at 0.9 and 4 h after stroke onset are presented in Figure 4. A slight decrease in ADC values was measured in the penumbra area, above the threshold for infarction but nevertheless indicating that some degree of cellular edema had occurred as a result of a perturbation of the ionic homeostasis. The intra-/extra-cellular compartmental volume shift resulting from this cell swelling may well explain this decrease in Na signal, recalling that the measured signal represents an average across all tissue compartment volumes and hence a reduced signal would accompany a decrease in the volume of the high-concentration extracellular compartment. A similar decrease of $11 \pm 8\%$ in Na signal was detected in a rabbit model in a region that ultimately infarcted, although it was not possible to say whether this measured decrease was in core or penumbra tissue [28]. Furthermore, a 13% decrease in Na signal was detected in a spectroscopic study of a canine model, but again no discrimination between core and penumbra was possible [44]. Nevertheless, if verified in further studies, this decrease in Na signal could help identify the presence of penumbra tissue in the acute phase, and hence a PWI-Na mismatch approach may be warranted, or indeed Na-MR alone [27]. A further consequence of these recent findings is the potential error which will be introduced into any stroke onset time determination, via extrapolation of the Na signal intensity back in time, in brain regions where the Na signal may have experienced a delay or even decrease before finally increasing; rather, such an analysis must be carried out exclusively in regions that infarcted immediately or very soon after onset, information that most likely will not be available in a clinical situation, and hence the usefulness of this approach is doubtful.

**Potential of Na-MRI in the acute phase**

From an MRI perspective, the currently available techniques of PWI and DWI have not been shown to offer a definitive identification of patients who will benefit from a thrombolytic treatment (some clinical trials have reported a benefit [45], while others have not [46]). Thresholds of blood flow (separating infarction from benign oligemia, for example) remain uncertain, while numerous studies have shown that ADC values...
The gradual increase in contralateral hemisphere (right side in the images) 0.9 sodium signal in confirmed penumbra tissue (dashed white line) is clearly visible at sodium signal in core tissue (solid white line) is evident. However, the decrease in sodium signal in confirmed penumbra tissue (dashed white line) is clearly visible at 0.9 h, which has partially (but not fully) recovered to normal levels by 4 h.

Scope for improvement
The quality of Na images have improved dramatically since the first human stroke study back in 1993 (see comparison in Figure 5), driven primarily by developments in magnet designs allowing for higher field strengths, RF coil design and ultra-short echo time sequences. While it is impossible to predict what lies ahead particularly in the brain. It can also be reasonably expected that developments in the wider MRI world will impact positively on Na imaging, for example novel image reconstruction methods for undersampled data such as iterative reconstruction [47] and compressed sensing [48], acquisition schemes such as RF encoding, as demonstrated recently in Na phantom experiments [49], or indeed improved RF pulse designs based on optimal control theory that may allow for the selective excitation of the intracellular component [50].

Many advances have been made in the design of ultra-short echo time imaging pulse sequences specifically aimed at Na-MRI, which have increased SNR and allowed for reduced acquisition times. The first such ultra-short echo time sequences developed were twisted projection imaging [36], followed in later years by 3D Cones [51], 3D projection-reconstruction [35] and most recently density-adapted 3D reconstruction [49].

While it is impossible to predict what lies ahead with any degree of certainty, one could reasonably expect further improvements particularly considering the rapid pace of developments in this field. For example, benefits may accrue from the use of phased array coil designs for Na-MRI particularly in the brain. It can also be reasonably expected that developments in the wider MRI world will impact positively on Na imaging, for example novel image reconstruction methods for undersampled data such as iterative reconstruction [47] and compressed sensing [48], acquisition schemes such as RF encoding, as demonstrated recently in Na phantom experiments [49], or indeed improved RF pulse designs based on optimal control theory that may allow for the selective excitation of the intracellular component [50].

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projection-reconstruction [31] and its improved 2D variant [52]. Other techniques have been successfully adapted for use in Na imaging, such as single point ramped imaging with T₁ enhancement (SPRITE), where a Centric single-point ramped imaging with T₁ enhancement variant was recently used to quantify the TSC in brain tumors [53], while the sweep imaging with Fourier transform (SWIFT) technique may prove useful due to its ability to acquire the entire signal from the short T₂ Na component due to its effective zero echo time [54]. This remains an active area of investigation and will hopefully lead to further innovations in the coming years.

Another interesting development in the field are efforts to introduce additional contrast into Na images via T₁/T₂ relaxation time or magnetization-prepared approaches. Currently, most studies are either spin-density weighted (that is, they measure the TSC) or they use a simple T₁-weighted approach (via the use of a short TR) in order to increase the SNR per unit scanning time via signal averaging, while other short TR/flip angle combinations have been used to minimize T₁-weighting with a view to optimizing the achievable SNR [55]. A recent study by Nagel et al. [56] compared these two conventional Na-MRI techniques with an inversion recovery-based approach (originally introduced in 2005 [57]) that was used to reduce the signal from edema. The resulting heavily T₁-weighted images exhibited a lower Na signal in the muscles of patients with chondropathies, which, it was hypothesized, may be due to a reduction in the T₁ relaxation times in the intracellular compared with the extracellular compartments. This inversion recovery-based preparation approach was also used to improve the quantification accuracy of the bound Na concentration in the extra-cellular space in cartilage by suppressing the free Na signal from fluid in and around the knee in osteoarthritic patients [13]. In another study, researchers were able to differentiate between brain tumor grades using a combination of spin-density- and relaxation-weighted Na sequences [58]. Measurements of the long T₁ components in the brain were reported as far back as 2004 [59], although efforts to accurately quantify the short T₂ component remain unsuccessful even at 7 T [30]. Nevertheless, T₂⁺ maps (from the long T₂ component) were recently obtained in the brain using a 3 T scanner, albeit with an acquisition time of 60 min [60]. In another study using a closely fitting surface transceiver coil, T₂⁺ times for both the short and long T₂ components (in addition to T₁-times) were quantified in the breast, illustrating that such measurements are indeed possible using optimized detector coils [63]. The possibility of introducing T₂ relaxation-time contrast into Na images may have particular relevance to stroke imaging, where the redistribution of Na⁺ between the intra- and extra-cellular compartments will be accompanied by a change in T₂ values, and hence a technique capable of introducing T₂-weighted contrast may well reveal more subtle effects that are presently obscured. Equally, knowledge of the changes that occur in T₁ values during the acute stroke period may also prove beneficial, if only
to avoid ambiguities arising from T1-saturation effects when pushing towards minimum TR values.

There are two other existing techniques that are capable of separating, to some degree, the intra- and extra-cellular compartment signals. The first involves the use of shift reagents such as TmDOTP(5-), first introduced in 2001 [62]. However, all such reagents are highly toxic to humans and it remains to be seen whether nontoxic variants can be developed. The second approach is the use of multiple quantum (MQ) coherences; in tissue, the quadrupolar interaction between the spin 3/2 moment of the Na nuclei and the local electric field gradients dominate and hence triple quantum (TQ) coherences are possible. Although signal from both intra- and extra-cellular Na ions exhibiting biexponential relaxation may pass through a TQ filter, the hope is that MQ techniques may be more selective for the intracellular component. In a nonhuman primate stroke study, the TQ Na signal was found to increase by 226 ± 70% within 0.9 ± 0.4 h of the stroke onset time, while little change was observed in the single quantum signal (n = 3) [63]. A spectroscopic study investigating the effect of glycemia levels on single and double quantum Na signals in the rat forebrain in a transient ischemia model concluded that double quantum signal intensity is a more sensitive monitor of changes in intracellular Na concentration during and after ischemia [64]. Thus, while there is no conclusive evidence to date that one can separate the intra- from the extra-cellular components using MQ techniques, nevertheless, the different evolutions in the single quantum versus MQ signal in the acute period observed in some studies warrants further investigation. The main drawback with MQ techniques is the drastically reduced SNR, which is approximately 10% of that obtainable in conventional single quantum coherence Na-MRI images, and hence this technique will benefit from an SNR boost from future technical developments. Hancu et al. succeeded in acquiring TQ Na images of the brain in 20 min using a 3 T scanner in 1999 [65], and while problems related to B0 inhomogeneities persist, several recent studies have made considerable progress in reducing their severity and increasing the SNR [66–68].

In stroke imaging, the question of whether or not to quantify the TSC remains open. On the one hand, accurate knowledge of the TSC would be essential if a threshold for tissue viability were to be determined and also, for example, in patients with bilateral stroke where relative measurements (contra- to ipsi-lateral) could be insensitive to any changes (or at least unreliable). However, TSC changes due to normal aging processes or indeed due to the presence of pathologies such as cognitive impairment (e.g., Alzheimer’s disease) arising perhaps due to cellular loss or shrinkage, are largely unknown (although a very recent study has looked at the effects of normal aging in the brain at 7 T [69]) and thus a relative comparison of the TSC in ipsi- versus contra-lateral regions may be warranted wherein the patient would act as their own control. Relative measures avoid some of the difficulties experienced with accurate TSC quantification and are certainly easier to implement in a clinical environment, although it can be expected that ongoing developments in the field, for example increasing the SNR, will make quantification more robust. Already studies have demonstrated that precise and accurate quantification of TSC in the brain can be obtained in rodents [26] and humans [1] with good spatial resolution in sub-10 min scans using commonly-available imaging platforms for each species (a 7 T research system and a clinical 3 T scanner, respectively). However, to push the spatial resolution even further to, say, an isotropic voxel of size 6 mm in humans (compared with 7.5 mm used in [1]) with accurate quantification using current coil hardware designs and imaging sequences currently requires higher field strength systems (7 T and above) [70], which are not widely available nor are they likely to be in the near future.

In conclusion, the field of Na-MRI has undergone a rapid pace of development particularly in the last decade and remains the focus of considerable research effort. It is clear that improvements in SNR will be required if Na-MRI is to play a central role in acute stroke imaging for several reasons: firstly, to detect the subtle signal changes that are likely to prove critical for penumbra tissue identification, whether via improved quantification accuracy or relative measurements, within a reasonable acquisition time (<10 min); and secondly, to allow for better spatial resolution in order to minimize partial volume effects arising, for example, when regions of cerebrospinal fluid are inadvertently included in the analysis where the threefold more Na concentration in cerebrospinal fluid compared with gray or white matter will skew the data. This problem is likely to be exacerbated in aged patients where brain atrophy has occurred. Nevertheless, given the improvement in image quality achieved thus...
far, one can remain optimistic for the future of the field.

**Future perspective**

It can be expected that the next 5–10 years will see the rapid pace of developments in the field of Na-MRI continue, with further improvements in the achievable SNR per unit time enabling accurate quantification of the TSC at better spatial resolutions in the human brain. The adoption of Na-MRI of new ideas in compressed sensing and iterative reconstruction coupled with developments in hardware and imaging pulse sequence developments are opening up many new possibilities, for example making it possible to accurately quantify the tissue sodium concentration (and perhaps also the intracellular compartment) and moving beyond mere Na spin-density-weighted imaging with the introduction of relaxation-time contrast or indeed Na relaxometry will yield further work in this area can be expected. Additional clinical studies involving acute stroke patients are required to inform on the usefulness of Na-MRI in this time-critical period, which will be aided by the growing availability of Na-MRI techniques.

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**Executive summary**

**Background**

- Considerable progress has been made in the field of sodium (Na)-MRI over the past decade, such that reasonable quality brain images can now be obtained using clinical 3 T scanners with 10-min acquisition times.

**Link between Na & stroke**

- From a stroke perspective, Na-MRI may offer a rapid, noninvasive method of identifying patients suitable for a therapeutic intervention.
- Changes in the intracellular Na concentration are known to occur shortly after an ischemic insult due to the impaired functioning of the Na⁺/K⁺ ATPase, which is responsible for maintaining cellular homeostasis.

**Clinical & preclinical stroke Na-MRI studies**

- Several studies in both animal models of stroke and in stroke patients have detected this change in Na imaging experiments, with early increases observed in core, infarcted tissue from an early time-point after stroke onset.
- Evidence in animal models suggests that the Na signal averaged across all tissue compartments may, in fact, decrease in penumbra tissue in the acute stroke phase, potentially offering a method of detecting penumbra tissue (i.e., underperfused tissue that is at risk of infarction but still amenable to salvage).

**Scope for improvement**

- Continuing improvements in image quality through hardware and imaging pulse sequence developments are opening up many new possibilities, for example making it possible to accurately quantify the tissue sodium concentration (and perhaps also the intracellular compartment) and moving beyond mere Na spin-density-weighted imaging with the introduction of relaxation-time contrast.

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Future Science Group
How can sodium MRI techniques help us understand acute stroke?


Fleischer L, Oesingmann N, Inglese M. B(0) inhomogeneity-insensitive triple-quantum-filtered sodium imaging using a 12-step phase-cycling scheme. NMR Biomed. 23(10), 1191–1198 (2010).


Provides an in-depth discussion of the true spatial resolution obtainable at present on 3T and 9.4 T platforms for quantitative TSC mapping.