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A. Hagiwara, M. Bydder, T.C. Oughourlian, J. Yao, N. Salamon, R. Jahan, J.P. Villablanca, D.R. Enzmann and B.M. Ellingson

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© A. Hagiwara, [®]M. Bydder, [®]T.C. Oughourlian, [®]J. Yao, [®]N. Salamon, [®]R. Jahan, [®]J.P. Villablanca, [®]D.R. Enzmann, and [®]B.M. Ellingson

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ABSTRACT

SUMMARY: Sodium MR imaging has the potential to complement routine proton MR imaging examinations with the goal of improving diagnosis, disease characterization, and clinical monitoring in neurologic diseases. In the past, the utility and exploration of sodium MR imaging as a valuable clinical tool have been limited due to the extremely low MR signal, but with recent improvements in imaging techniques and hardware, sodium MR imaging is on the verge of becoming clinically realistic for conditions that include brain tumors, ischemic stroke, and epilepsy. In this review, we briefly describe the fundamental physics of sodium MR imaging tailored to the neuroradiologist, focusing on the basics necessary to understand factors that play into making sodium MR imaging feasible for clinical settings and describing current controversies in the field. We will also discuss the current state of the field and the potential future clinical uses of sodium MR imaging in the diagnosis, phenotyping, and therapeutic monitoring in neurologic diseases.

ABBREVIATIONS: ESC = extracellular sodium concentration; IDH = isocitrate dehydrogenase; ISC = intracellular sodium concentration; NHEI = Na+/H+ exchanger isoform 1; TSC = total sodium concentration

Routine clinical MR imaging is performed exclusively by using hydrogen nuclei (ie, protons, H⁺) because it is the most abundant element in the human body in the form of water. However, other nuclei are also detectable with MR imaging and may provide complementary physiologic information to conventional proton MR imaging. Referred to collectively as "X-nuclei," elements including sodium (²³Na), potassium (³⁵K), chloride (³⁵Cl), and phosphorus (³¹P) all have detectable magnetic moments and all play critical roles in the biochemistry of living tissues.

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From the UCLA Brain Tumor Imaging Laboratory (A.H., M.B., T.C.O., J.Y., B.M.E.), Center for Computer Vision and Imaging Biomarkers, and Department of Bioengineering (J.Y., B.M.E.), Henry Samueli School of Engineering and Applied Science, University of California, Los Angeles, Los Angeles, California; and Department of Radiological Sciences (A.H., M.B., J.Y., N.S., R.J., J.P.V., D.R.E., B.M.E.), Neuroscience Interdepartmental Program (T.C.O., B.M.E.), and Department of Psychiatry and Biobehavioral Sciences (B.M.E.), David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, California.

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Please address correspondence to Benjamin M. Ellingson, PhD, UCLA Brain Tumor Imaging Laboratory, Departments of Radiology and Psychiatry and Radiological Sciences and Psychiatry, David Geffen School of Medicine, University of California, Los Angeles, 924 Westwood Blvd, Suite 615, Los Angeles, CA 90024; e-mail: bellingson@mednet.ucla.edu

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Sodium is the second most abundant element in the body detectable on MR imaging. Sodium homeostasis is crucial for life because it is a major determinant of body fluid osmolality, and sodium sensing is performed in the brain by specialized sodium channels within the circumventricular organs to maintain a range of 135–145 mM. Sodium also plays a crucial role in the propagation of neural signals in and between neurons, and disruption in sodium homeostasis as well as structural and metabolic integrity has been identified in a variety of neurologic disorders including brain tumors, stroke, and epilepsy. Hence, sodium MR imaging has remarkable potential for use in diagnosis, characterization, and treatment monitoring in neurologic diseases.

The fundamental limitation to translational use of sodium MR imaging for clinical care is the low inherent MR signal. The MR imaging signal is approximately 10,000 times lower than that of protons, which is due to a combination of the following: 1) a concentration of about 0.055% of that of water protons; 2) a gyromagnetic ratio (γ , relates the main magnetic field strength, B₀, to the resonance frequency) of about 26.5% of that of the proton, thus a lower energy ($\Delta E = \gamma \hbar B_0$) and lower inherent bulk magnetization; and 3) a nuclear spin of 3/2, leading to electric quadrupolar interactions between the nucleus and its environment resulting in faster T2* decay. Overcoming these issues and achieving adequate MR signal levels necessitates acquisition schemes with low spatial resolution, short TE, multiple measurements, and long scan times.

Researchers have put considerable effort into optimizing these various factors and taking advantage of both the acquisition algorithm and hardware improvements. In this review, we briefly describe the fundamental physics of sodium MR imaging tailored to the neuroradiologist, focusing on the basics necessary to understand factors that are involved in making sodium MR imaging feasible for clinical settings and describing current controversies in the field, including the measurement of intracellular sodium concentration. We will also discuss the current state of the field and the potential future clinical uses of sodium MR imaging in the diagnosis, phenotyping, and therapeutic monitoring in neurologic diseases.

Sodium MR Imaging Physics and Acquisition

In practice, MR imaging of sodium ions is essentially the same as imaging protons, but with lower SNR and shorter (and more complex) T2 relaxation characteristics. Due to the inherent nuclear spin of 3/2, the sodium ion exhibits electric quadrupolar interactions between the nucleus and its environment, 7,8 leading to what appears to be a biexponential T2 decay in many biologic tissues. The short T2 species appears to occur most often in viscous liquids or semisolid tissues, where there is a continuum of electric field gradients (depending on the orientation of the sodium ion and the position of an anion) that produces a broad range of energy levels and gives rise to short T2 relaxation. A longer T2 component is observed from the energy levels unperturbed by the electric field gradients, such as those in CSF and other nonviscous liquids because the motion of the ions causes electric field gradients to average to zero over the measurement time scale, resulting in a single, longer T2 relaxation. This observation has been replicated in the human brain, where parenchyma, including both white and gray matter, exhibits a characteristic biexponential T2 decay, while the nonviscous CSF displays monoexponential T2 relaxation.9

Most measurements used in clinical sodium MR imaging focus on estimation of the total sodium concentration (TSC) or the volume-weighted average of the intra- and extracellular sodium concentrations (ISC and ESC, respectively). Notably, ISC and ESC do not correspond to short and long sodium T2 relaxation (explained later in this section). Because ESC is stable at around 140 mM, TSC is mainly affected by changes in ISC and alterations in the volume fractions between the intra- and extracellular space. TSC in the brain was reported to range from 30 to 56 mM¹⁰⁻¹² and remain constant throughout adulthood in cognitively healthy individuals, and its regional variation in the brain was reported to be minimal.¹² Sodium concentration in the CSF has been known to show circadian fluctuation; 13 however, whether TSC in the brain tissue also shows circadian fluctuation and whether the degree of fluctuation is different between normal and pathologic brain tissues are unknown. Remarkable effort has been made to measure ISC during the past decades in animal models to elucidate cellular homeostasis, energetic state, and functionality of sodium pumps; however, these measurements involve the use of highly toxic extracellular paramagnetic "frequency shift reagents" that are trapped within the extracellular space and shift the resonance frequency of extracellular sodium,

resulting in isolation of purely intracellular sodium ions. ^{14,15} Unfortunately, due to their toxicity and the fact they do not cross the blood-brain barrier, shift reagents for sodium MR imaging are not used in humans.

Two major physics approaches to measure the ISC in humans that have been explored include the use of inversion recovery 16-19 and multiple quantum filtering 20-22 techniques. Similar to FLAIR, the inversion recovery approach assumes that a simple inversion pulse can suppress the sodium signals free from macromolecules; hence, the sodium signal in the extracellular space is also presumed to be suppressed. However, there is evidence that the T1 relaxation times of intra- and extracellular sodium are very similar, 23-28 and there is little reason to assume that the T1 of the sodium within the extracellular compartment is more similar to that in CSF than in the intracellular compartment. Indeed, extracellular fluid in brain tissue is largely different from CSF in composition. 29

Multiple quantum filtering is a relatively sophisticated approach that works on the premise that the slow-moving sodium, presumably in the intracellular space, can be selectively detected on the basis of the underlying biexponential T2 relaxation of sodium.³⁰ However, molecular interactions that result in equivalently small T2 values in both intra- and extracellular compartments are expected to result in similar multiple quantum coherences.²⁷ Indeed, previous experiments have shown a contribution of the extracellular sodium to the multiple quantum coherences detected by multiple quantum-filtering sodium MR imaging, and the degree of contamination is largely unexplored.³¹⁻³³ Therefore, direct measurement of the ISC solely using sodium MR imaging physics without the use of exogenous contrast agents is not yet plausible, and such claims in previous literature using the relaxation behavior of sodium should be cautiously interpreted. Furthermore, the exchange rate of sodium between intra- and extracellular space, which might affect the measurement of the ISC, has not been considered in previous studies. Future studies aimed at using multiparametric input from proton diffusion MR imaging and/or PET may be useful for using estimates of cell density³⁴ to disentangle the ISC from the TSC measured by sodium MR imaging, but as of now, TSC is the most reliable measurement parameter for routine sodium MR imaging examinations.

The primary challenge for clinical sodium MR imaging is the very short T2, meaning that most of the sodium MR imaging signal is lost within a few milliseconds. Hence, imaging with a very short, ultrashort, or zero TE (ie, TE of <1 ms) is almost mandatory for any sodium MR imaging application. High quantification accuracy of the sodium concentration with ultrashort TE imaging has been reported. 10,35 To reduce the duration of signal readout while maintaining sufficient SNR, non-Cartesian sequences (eg, radial or spiral trajectories) are preferred. 36,37 For example, Ridley et al³⁸ used a 3D radial projection for whole-brain imaging, with 3-mm isotropic resolution on a 3T clinical scanner in 34 minutes. On the other hand, Thulborn et al³⁹ showed that a twisted radial k-space imaging trajectory could be used for whole-brain coverage with 5-mm isotropic resolution (44 slices) on a 3T clinical scanner in approximately 8 minutes, which was sufficient for brain tumor imaging applications.

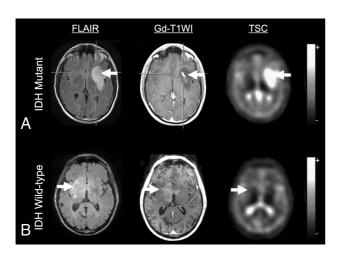


FIG 1. A, MR imaging of a 38-year-old male patient with an *IDH*-mutated glioblastoma, World Health Organization grade IV. The tumor (*white arrows*) shows focal contrast enhancement in the TI-weighted image and is clearly depicted in the FLAIR image. Sodium imaging shows increased TSC. *B*, MR imaging of a 78-year-old male patient with an *IDH* wild-type anaplastic astrocytoma, World Health Organization grade III, in the right basal ganglia. The tumor (*white arrows*) shows focal contrast enhancement in the TI-weighted image and diffuse abnormalities in the FLAIR image. Sodium imaging shows no abnormality. Adapted and reproduced with permission from Shymanskaya et al.²²

In addition to creative k-space trajectories, advancements in hardware have also made sodium MR imaging more clinically realizable. The use of high- and ultra-high-field strength scanners, for example, proportionally increases the SNR, making commercial 7T MR imaging scanners feasible for faster or higher resolution sodium imaging.¹¹ Of note, higher resolution, especially with 3D imaging, helps reduce partial volume effects. In addition to the use of high-field-strength scanners, receiver coil architecture is also of great importance because the design of this coil predetermines the maximum attainable SNR. Because proton MR imaging is often performed concurrently with sodium imaging, the same physical coil housing (ie, "dual tuned" coils) is desired. Dual-tuned designs of different Larmor frequencies come with unique challenges, however, such as coupling between the sodium and proton coils. 40 This issue could be addressed by using 2-coil geometries that are intrinsically decoupled, dual-tuning a single coil, or strategies to actively decouple the 2 coils.⁴⁰ Use of array coils is an attractive approach in sodium imaging to further increase the SNR and reduce scan time by using parallel imaging strategies. 41 For example, Lee et al 42 demonstrated up to a 400% increases in the SNR using a 4-channel coil >20 years ago. To date, the highest number of channels reported for a head sodium coil is 32,²⁸ while 20- to 30-channel coils are commercialized and available through third-party coil vendors. However, despite remarkable progress, array coils are still rarely used in sodium imaging because of hardware and software limitations and additional costs. 41 For using array coils in sodium MR imaging, the receivers should be capable of handling the frequencies relevant to sodium acquired with analog-to-digital converters and having processing engines capable of sorting and combining the signals from each coil properly. Most commercial scanners

have offered only a single broadband X-nuclei receiver channel, limiting the use of array coils.

Neurologic Applications

Brain Tumors. In brain tumors, sodium MR imaging has the potential to reveal molecular information related to cell viability, proliferation, migration, invasion, and immunogenicity⁴³⁻⁴⁵ and may enable us to reveal molecular responses to treatment before morphologic changes can be observed. TSC is reportedly elevated in brain tumors both in humans 19,22,39,46-48 and animals, 49,50 a feature that may be due to the ISC, the volume of the extracellular space, or both, considering that the ESC remains relatively constant and much higher than the ISC as long as there is moderate tissue perfusion.⁵¹ Increases of the ISC in tumors are partly related to the increased energy demand arising from cell proliferation, because negative sodium gradients across the cell membranes are maintained by consumption of adenosine triphosphate.⁵² In addition, the Na⁺/H⁺ exchanger isoform (NHE1) (SLC9A1) is upregulated in gliomas and is a potential therapeutic target due to its role in the progression of malignant gliomas, 53-55 influence on pH homeostasis in glioma cells,54-57 influence on seizure activity,6 and potential increased resistance to both chemoradiation⁵⁸ and anti-PD-1 immunotherapy.⁵⁶

Although the literature is sparse, studies have shown differences in sodium MR imaging contrast among different tumor grades^{48,59} and between active tumors and peritumoral edema⁴⁶ or other types of lesions, 60 and these differences appear to reflect the general prognosis, ⁵⁹ with higher TSC in areas of more aggressive tumor. Despite this general trend, a number of studies have shown differences in sodium MR imaging contrast between isocitrate dehydrogenase (IDH) mutant and wild-type gliomas, 22,59,61 with IDH-mutant human gliomas showing higher TSC than IDH wild-type gliomas (Fig 1).^{22,61} While this finding is counterintuitive, it could be due to the lower cellular density of IDH-mutant gliomas,⁶² leading to a larger extracellular space and higher TSC. Alternatively, IDH-mutant gliomas often result in more frequent seizures compared with more aggressive high-grade IDH wildtype malignant gliomas, 63,64 and because expression of NHE1s is strongly linked to seizure activity,6 this finding may also explain the differences in TSC observed between IDH-mutant and wildtype gliomas.

Sodium MR imaging has also shown some promise in identifying the early treatment response in brain tumors. 39,65,66 For example, in rat glioma models, response to chemotherapy using sodium MR imaging was detectable even earlier than in proton diffusion MR imaging.⁶⁷ In rat models of subcutaneously implanted gliomas, lower total sodium signals were observed in gliomas treated with 1,3-Bis(2-chloroethyl)-1-nitrosourea compared with untreated gliomas.⁶⁸ Another study showed a pronounced increase in the TSC following 1,3-Bis(2-chloroethyl)-1nitrosourea treatment in orthotropic rat gliomas compared with untreated gliomas.⁵⁰ Notably, the increase in TSC after treatment occurred before tumor shrinkage. Even though the discrepancy in the results between these 2 studies may partly lie in differences in the implantation sites of gliomas and acquisition methods, the increase in TSC observed in the latter study may also be due to a combination of treatment response leading to necrosis and an

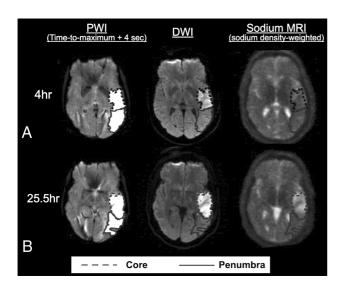


FIG 2. Images of a representative section from a patient with ischemic stroke showing the hypoperfused (time-to-maximum + 4 seconds) perfusion maps, the DWI with a DWI-hyperintense core in the *dotted outline*, the PWI-DWI mismatch tissue (penumbra) in the *solid outline*, and sodium images for (*A*) 4 and (*B*) 25.5 hours after the onset. This patient had a perfusion/diffusion mismatch at the first time point. The absolute lesion volume of the core enlarged from the first to the second time point, while the penumbral volume diminished. Note that the sodium signal is not increased in the first time point, while the high sodium signal is matched with DWI hyperintensity at the second time point. Adapted and reproduced with permission from Tsang et al. ⁷⁶

increase in the extracellular space. Additionally, Thulborn et al³⁹ evaluated the effects of the standard chemoradiotherapy on 20 patients with human glioblastomas using sodium MR imaging and noted an increase in TSC after successful treatment.

Acute Ischemic Stroke. Acute ischemic stroke occurs due to sudden occlusion of arteries within the brain, resulting in reductions of adenosine triphosphate production and Na⁺/K⁺-ATPase activity. Inadequate Na+/K+-ATPase activity disrupts the ion homeostasis, leading to an increase in the ISC, cytotoxic cell swelling, and eventual cell death.⁶⁹ Sodium MR imaging may be useful as a surrogate marker of Na⁺/K⁺-ATPase activity and cell viability in the ischemic tissue, with potential implications for determining tissue viability. 70 Monotonic increases in the TSC after acute ischemic stroke have been reported both in animals 71,72 and humans, 73,74 using time scales relevant for patient management (ie, 0-24 hours following onset), and this increase does not appear to normalize in the natural course following stroke. In contrast, decreased ADC in the acute phase normalizes when vasogenic edema starts in the subacute phase (ie, 24-72 hours following onset).⁷⁵ Hussain et al⁷³ demonstrated that there was a 10% increase in sodium signal in the first 7 hours, followed by a rapid increase in sodium until a plateau of a 69% increase at 48 hours relative to baseline values, during which time the ADC did not fluctuate. Because sodium concentration correlates with the duration of ischemia, the onset time may be more accurately estimated by the sodium concentration than diffusion MR imaging changes, providing potential utility in "wake-up" strokes.

Additional studies have demonstrated that sodium MR imaging signals in the region with a perfusion-diffusion mismatch may not differ from those in contralateral normal tissue until around 32 hours after symptom onset, indicating that sodium MR imaging may help identify the viable tissue in the penumbra, even when the onset time of a stroke is unknown (Fig 2).⁷⁶ Despite these initial studies, the specific thresholds of TSC for determining reversible and irreversible ischemic tissues and the vulnerability of infarcted tissues to hemorrhage following reperfusion therapy are yet to be determined. Conceivably, sodium MR imaging in combination with conventional imaging techniques may enable more judicious selection of candidates for endovascular thrombectomy in the future, rather than using a fixed time window as is the current practice.

Epilepsy. Because sodium homeostasis affects neuronal excitability, sodium MR imaging has the potential to detect subtle disturbances in sodium concentration in seizure disorders including epilepsy. Several pathologic mechanisms in epilepsy are implicated in the change of the TSC observed in the brain, with the primary mechanism being dysfunction of sodium channels and Na⁺/K⁺-ATPase in patients with epilepsy due to genetic mutation and mitochondrial dysfunction, leading to depolarization and the increase in the ISC. Tr-79 Additionally, reduction in the size of the extracellular space due to an increase in the intracellular osmolarity can occur during seizure activity, so,81 and an increase in the extracellular space due to neuronal loss, gliosis, and blood-brain barrier disruption, arising from chronic epilepsy or underlying diseases such as stroke or trauma, can lead to alterations in TSC measurements using sodium MR imaging. 22

Despite the potential impact, to date, very few sodium MR imaging studies have been performed in epilepsy and seizure disorders. Wang et al⁸³ reported a dynamic postictal temporal change in proton ADC and TSC using sodium MR imaging after the administration of kainic acid to rats. In the pyriform cortex and amygdala, decreases in the ADC were noted as early as 5 hours after kainic acid administration, and the ADC values further decreased until 24 hours after the seizures. ADC values returned to normal levels 7 days postictally. Meanwhile, the TSC did not change at 5 hours postictally but increased at 24 hours and remained elevated even at 7 days postictally. These changes in ADC and TSC were interpreted as being influenced by sodium entry into the excited neurons and accompanying cellular swelling, followed by energy deficiencies and cell death, in line with pathologically-confirmed extensive neuronal cell loss by day 7. However, future research is desired to elucidate the relationship between the change in sodium concentration and pathophysiology in more detail by using shift reagents in animal models, to determine the degree of contribution by the change in the ISC to the increased TSC.

Additionally, Ridley et al³⁸ used sodium MR imaging to examine 9 patients with epilepsy during interictal periods and 1 patient who incidentally presented with several seizures during the MR imaging examination. TSC in the intracerebral electroencephalogram-defined epileptogenic regions was increased in the interictal group, a finding that can be explained by an increase in the ISC due to voltage-gated sodium channel mutations leading to a persistent

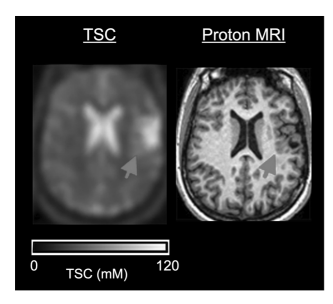


FIG 3. A patient with a cortical malformation. The TSC map shows high value on a cortical malformation that is subtle on high-resolution proton 3D-TI-weighted image (*arrows*) (adapted and reproduced with permission from Ridley et al³⁸). However, the effect of the partial volume effect should be examined in a future study with the achievement of higher-resolution sodium MR imaging.

inward sodium current, along with an increase in the extracellular space due to cell loss and glial formation (Fig 3).^{6,82} In contrast to patients in the interictal period and consistent with preclinical studies, the TSC was slightly decreased in the epileptogenic area in the patient who presented with multiple seizures during the MR imaging examination. Together, these preliminary studies suggest that sodium MR imaging may be useful for identification and illumination of epileptic activity, but important questions remain, including the precise temporal changes in TSC that occur during and after seizure activity, the effects of antiepileptic medication, the sensitivity of sodium MR imaging for epileptic foci detection compared with standard proton MR imaging, and the value of sodium MR imaging as a tool to predict surgical outcome in patients with refractory epilepsy.

CONCLUSIONS

Sodium MR imaging has the potential to complement routine proton MR imaging examinations with the goal of improving diagnosis, disease characterization, and clinical monitoring in neurologic diseases. In the past, the utility and exploration of sodium MR imaging as a valuable clinical tool have been limited due to the extremely low MR signal, but with recent improvements in imaging techniques and hardware, sodium MR imaging is on the verge of becoming clinically feasible for conditions including brain tumors, stroke, and epilepsy.

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