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Sodium Magnetic Resonance Imaging of the Human Heart at 7.0 T: Preliminary Results

Andreas Graessl 1, Anjuli Ruehle 1, Helmar Waiczies 3, Ana Resetar 2, Stefan H. Hoffmann 2, Jan Rieger 3, Friedrich Wetterling 4, Lukas Winter 1, Armin M. Nagel 2 and Thoralf Niendorf 1,5

1 Berlin Ultrahigh Field Facility (B.U.F.F.), Max-Delbrueck-Center for Molecular Medicine, Berlin, Germany
2 Division of Medical Physics in Radiology, German Cancer Research Center (DKFZ), Heidelberg, Germany
3 MRI.TOOLS GmbH, Berlin, Germany
4 Trinity College, Institute of Neuroscience, The University of Dublin, Ireland
5 Experimental and Clinical Research Center, A joint cooperation between the Charité Medical Faculty and the Max-Delbrück Center for Molecular Medicine, Berlin, Germany

Corresponding Author: Prof. Dr. Thoralf Niendorf
Berlin Ultrahigh Field Facility (B.U.F.F.)
Max-Delbrueck-Center for Molecular Medicine
Robert-Roessle-Strasse 10
13125 Berlin
Germany
Tel.: +49 30 9406 4505
Fax.: +49 30 9406 49178
E-Mail: Thoralf.Niendorf@mdc-berlin.de
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Abstract

Objective: Examine the feasibility of three-dimensional and whole heart coverage \(^{23}\text{Na}\) cardiac MRI at 7.0 T including single cardiac phase and CINE regimes.

Methods: A four channel transceiver RF coil array tailored for \(^{23}\text{Na}\) MRI of the heart at 7.0 T (\(f=78.5\) MHz) is proposed. An integrated bow tie antenna building block is used for \(^1\text{H}\) MR to support shimming, localization and planning in a clinical workflow. Signal absorption rate simulations and assessment of RF power deposition were performed to meet the RF safety requirements. \(^{23}\text{Na}\) cardiac MR was conducted in an in vivo feasibility study. 3D gradient echo (GRE) imaging in conjunction with Cartesian phase encoding (total acquisition time \(T_{AQ}=6:16\) min) and whole heart coverage imaging employing a density-adapted 3D radial acquisition technique (\(T_{AQ}=18:20\) min) were used.

Results: For 3D GRE based \(^{23}\text{Na}\) MRI, acquisition of standard views of the heart using a nominal in-plane resolution of (0.5 x 0.5) \(\text{mm}^2\) and a slice thickness of 15 mm were feasible. For whole heart coverage 3D density-adapted radial \(^{23}\text{Na}\) acquisitions a nominal isotropic spatial resolution of 6 mm was accomplished. This improvement versus 3D conventional GRE acquisitions reduced partial volume effects along the slice direction and enabled retrospective image reconstruction of standard or arbitrary views of the heart. Sodium CINE imaging capabilities were achieved with the proposed RF coil configuration in conjunction with 3D radial acquisitions and cardiac gating. Cardiac gated reconstruction provided an enhancement in blood-myocardium contrast of 20% versus the same data being reconstructed without cardiac gating.
**Conclusions:** The proposed transceiver array enables $^{23}$Na MR of the human heart at 7.0 T within clinical acceptable scan times. This capability is in positive alignment with the needs of explorations that are designed to examine the potential of $^{23}$Na MRI for the assessment of cardiovascular and metabolic diseases.

**Keywords:**
magnetic resonance imaging; ultrahigh field MR; cardiovascular MRI; sodium imaging; RF coil technology; transceiver array
List of Abbreviations

$^{23}$Na MRI  sodium magnetic resonance imaging
$B_0$       main magnetic field strengths
$B_1^+$     electromagnetic transmission field
BMI         body mass index
CP          circular polarized
CNR         contrast-to-noise ratio
ECG         electrocardiogram
Na$^+/K^+$ ATPase sodium-potassium adenosine triphosphatase
EMF         electromagnetic fields
f           frequency in Hertz
FA          flip angle in degree
FLASH       Fast Low Angle Shot
FOV         field of view
FR4         copper clad sheet for electronic applications using glass-epoxy resin
IEC         International Electrical Commission
NA          number of averages
NEMA        National Electrical Manufacturers Association
Q$_{un}$    unloaded quality factor
Q$_l$       loaded quality factor
RF          radio frequency
RX          reception
S           scattering parameter
SAR         specific absorption rate
SAR$_{10g}$ specific absorption rate averaged over 10 g
SD          standard deviation
SNR         signal-to-noise ratio
T$_1$       longitudinal relaxation time
T$_2^*$     effective transversal relaxation time
TE          echo time
TR          repetition time
TX          transmission
UHF-MR      ultrahigh field magnetic resonance
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This pilot study demonstrated the feasibility of sodium MRI of the human heart at 7.0T using a dedicated transceiver RF coil array. The proposed setup afforded the acquisition of sodium images with reasonable myocardial signal in clinically acceptable scan times. 3D density adapted radial acquisitions yielded a signal gain compared to Cartesian gradient echo acquisitions. This improvement supported whole heart, CINE sodium imaging of the heart with an isotropic spatial resolution of 6 mm within approximately 19 minutes scan time.
Introduction

Knowledge about compromised cell integrity of myocardial tissue provides valuable information for the assessment of myocardial viability, with tissue sodium concentration being a potent indicator of impaired cellular ion homeostasis and cell membrane integrity. In healthy tissue the Na⁺/K⁺ATP-ase maintains a transmembrane concentration gradient with an intracellular sodium concentration of about 15 mmol/L against an extracellular concentration of about 140 mmol/L (1,2). During ischemia, for instance, the function of the sodium/potassium pump is compromised by declining energy reserves resulting in increased intracellular sodium concentration.

Sodium MRI (²³Na MRI) is a viable approach for gaining better insights into cellular metabolism with a broad spectrum of biomedical imaging research applications (3). Previous studies eloquently reported credible data on ²³Na MRI of the heart and demonstrated that ²³Na-MRI is suitable for the detection and assessment of acute and chronic heart disease due to increased sodium concentration after myocardial infarction (4-9). These pioneering explorations were primarily limited to animal studies. Constraints dictated by the rapidly decaying ²³Na signal and the low sensitivity of ²³Na MRI versus clinical ¹H-MRI induced scan times being not clinically acceptable and hence constituted a challenge for ²³Na MRI of the human heart. Notwithstanding this obstacle ECG gated ²³Na MRI of the heart was accomplished for magnetic field strengths commonly used in today’s clinical practice (B₀ ≤ 3.0 T) (10-14). Yet, the low gyromagnetic ratio of ²³Na and the low Na⁺ tissue concentrations compared with hydrogen resulted in long acquisition times and modest spatial resolution for ²³Na MRI.
of the heart. Although ample signal-to-noise was demonstrated for ventricular blood, the limited myocardial $^{23}$Na signal constitutes a challenge for cardiac sodium MRI en route to a clinical tool. Recognizing the sensitivity gain intrinsic to ultrahigh magnetic fields ($B_0 \geq 7.0$ Tesla) and yet unimpaired transmission field homogeneity (15) due to the comparably low $^{23}$Na resonance frequency which is close to $^1$H MRI at 1.5 T it is conceptually appealing to pursue cardiac $^{23}$Na-MRI at 7.0 Tesla.

Realizing the challenges of human $^{23}$Na MRI alongside the opportunities of ultrahigh field MR (UHF-MR), this study examines the feasibility of cardiac $^{23}$Na MRI at clinically acceptable acquisition times using free breathing 3D Cartesian gradient echo and cardiac gated 3D density-adapted radial acquisition techniques at 7.0 T. To meet this goal a local four-element transceiver RF surface coil that is customized for cardiac $^{23}$Na MRI at 7.0 T is proposed. To meet the safety and RF power deposition limit requirements of MRI, electromagnetic field (EMF) simulations and specific absorption rate (SAR) considerations are conducted. The feasibility and applicability of the proposed approach for $^{23}$Na MRI of the heart at 7.0 T is presented and its suitability for single cardiac phase and for cinematic $^{23}$Na MRI is demonstrated in initial volunteer studies, as a precursor to broader clinical studies. The merits and limitations of the proposed transceiver surface coil are discussed and implications for clinical MRI of the human heart at 7.0 T are considered.
Experimental

RF coil design

The proposed dual-frequency RF coil setup (Figure 1a) consists of separate elements for $^{23}$Na and $^1$H MR. For $^{23}$Na MR a four channel RF coil array was constructed using two modestly curved lightweight formers to conform to an average anterior and superior chest as illustrated in Figure 1a. Each section contains two rectangular loop elements (size: $\text{(210 x 140) mm}^2$) which each share a common conductor as shown in Figure 1b, c. The structure shown in Figure 1b was etched from 32 $\mu$m copper on 1 mm FR4 substrate. The conductor thickness was chosen to be more than four times the skin depth of 7.45 $\mu$m for copper at 78.6 MHz. Each loop element is divided into five sections by chip capacitors to form a uniform and balanced current distribution along the loop. Distributing the overall capacitance also reduces the voltage at every capacitor and thus reduces coupling to the sample as well as local SAR. A trimmer capacitor (C4) was integrated in the shared conductor to afford adjustable decoupling of the neighboring loop elements (16). The common conductor of the anterior coil section also includes a parallel resonant trap circuit (C6/L6) tuned to 297 MHz to prevent $^1$H signals to couple into the loop structure. The ports were connected to the coaxial RF feeding cables with a tune/match network consisting of a parallel tuning capacitor and a serial matching trimmer capacitor for each loop element. The coil was tuned to 78.6 MHz which corresponds to the resonance frequency of $^{23}$Na at 7.0 T and matched to 50 $\Omega$. Unbalanced currents on the coaxial cables were suppressed by
parallel resonant cable traps. Cable traps were designed as double-turn solenoids of the coaxial cable with an appropriate capacitor soldered to the outer conductor at the crossing of the cable.

Basic $^1$H imaging capabilities were achieved by a dipole bow tie element tuned to 297 MHz (Figure 1d) as described in (17). The dipole bow tie element was placed inside the left loop element of the anterior section of the $^{23}$Na coil as depicted in Figure 1 b. This approach provides basic means for localization, $B_0$ shimming and slice positioning based on $^1$H images.

Bench measurements and characterization of the RF coil performance were performed using an 8-channel vector network analyzer (ZVT 8, Rohde & Schwarz, Memmingen, Germany). The quality factor ratio $Q_U/Q_L$ of the loop elements was determined based on the measured resonance bandwidth with the array placed on an exemplary human subject. A high value indicates a predominant impact of sample noise over coil noise, which is a key requirement for the applicability of RF coil arrays.

**MR hardware**

MR experiments were conducted on a 7.0 T whole body MR scanner (Magnetom, Siemens Healthcare, Erlangen, Germany), equipped with an gradient system that offers a maximum slew rate of 200 mT/m/ms and a maximum gradient strength of 40 mT/m (Siemens Medical Solutions, Erlangen, Germany).

An 8 kW single channel RF amplifier was used for $^1$H MR. An 8 kW broadband single channel x-nuclei RF amplifier (Dressler HF-Technik GmbH, Germany) was used for $^{23}$Na MR. The x-nuclei amplifier output was split into 4 equal-amplitude signals by
means of three hybrid couplers (Anaren Microwave Inc., New York, US) compiled to a splitting cascade from 1:2 and 2:4. Phase adjustments of the single channels were implemented by phase-shifting coaxial cables connected to the power splitting network. The four $^{23}$Na elements and the one $^1$H element were connected to the RF system using an interface box with transmit/receive switches and integrated low-noise preamplifiers designed for the corresponding frequencies (Stark Contrast, Erlangen, Germany).

**Electromagnetic field simulations**

EMF and SAR simulations were performed using the finite integration technique of CST Studio Suite 2012 (CST AG, Darmstadt, Germany) together with human voxel model Duke (BMI: 23.1) (18). The simulations included the RF shield of the gradient coil. The voxel model was truncated at the neck and at the hips as displayed in Figure 1e, allowing for a high resolution mesh in the target region formed around the heart. Two simulations were performed with voxel model tissue parameters corresponding to 78.6 MHz and 297 MHz as listed in (19) to estimate SAR values for the respective frequencies. Decoupling capacitors and the feeding points of the elements were modeled as 50 Ω ports. Final field results were accomplished incorporating lumped decoupling, tuning and matching capacitors in the built-in circuit simulator of CST Studio Suite (CST Design Studio 2012), following the circuit co-simulation workflow proposed in (20). The capacitor values were optimized with respect to the S-parameter simulation and were used as a starting point for the practical realization. For the final configuration and phase settings used in in vivo measurements SAR values were calculated. The input power was adjusted to meet the regulations of the IEC guideline IEC 60601-2-33.
Ed.3 (21). At 78.6 MHz, which is in the frequency range of 1.5T proton imaging, transmission field non-uniformities are expected to present no major practical obstacle for $^{23}$Na MRI at 7.0 T. A circular polarized (CP) like mode is used for transmission. The CP mode refers to a phase setting with each channel driven by a phase corresponding to its angular position relative to the body center in the transversal plane (Channel1: 0°, channel2: -305°, channel3: -136°, channel4: -93°).

**Ethics statement**

For the in vivo feasibility study, subjects without any known history of cardiac disease were included after due approval by the local ethical committee (registration number DE/CA73/5550/09, Landesamt für Arbeitsschutz, Gesundheitsschutz und technische Sicherheit, Berlin, Germany). Informed written consent was obtained from each volunteer prior to the study.

**Cardiac imaging protocol**

In vivo cardiac MR was performed in three healthy subjects (2 male, 1 female; age range: 26 - 29; BMI range: 19 - 22, heart rate range: 62 - 78 bpm).

$^1$H imaging was performed using 2D CINE FLASH imaging (matrix size 256x256, TE=1.84 ms, TR=4.14 ms, voxel size (1.4x1.4x4.0) mm$^3$, number of cardiac phases=30, total acquisition time= 0:16 min). For cardiac imaging slice positioning was carried out following international consensus (22,23). For this purpose the heart was localised in three orthogonal thoracic slices placed along each main axis of the upper torso using
single breath-hold, low spatial resolution 2D gradient echo scout images. The long axis of the left ventricle was dissected twice, and finally a stack of short axes was obtained. These slices provided the basis for planning standard long axis views (four-chamber, three-chamber and two-chamber view) derived from 2D CINE FLASH imaging. Based on the four-chamber view, a mid-ventricular short axes view positioned parallel to the mitral valve plane was planned. The obtained slice positions were subsequently used for slab and 3D volume positioning for $^{23}$Na imaging.

For $^{23}$Na MRI of the heart flip angle calibration was done offline (Matlab, Mathworks, Natick, USA) using a curve fitting algorithm on a set of signal intensity points which were acquired with a series of RF transmitter reference amplitudes. For flip angle calibration the whole field of view was taken into account. $^{23}$Na MRI of the heart was performed with two imaging protocols:

- short axis views and four chamber views of the heart were acquired using untriggered, free breathing 3D gradient echo (3D FLASH) imaging with Cartesian phase encoding: matrix size 64 x 48, echo time TE=1.91 ms, repetition time TR=28 ms, voxel size (5 x 5 x 15) mm$^3$, number of averages 35, receiver bandwidth=120 Hz/pixel, flip angle= 28°, 8 slices per slab, total acquisition time 6:16 min.
- whole heart coverage $^{23}$Na datasets were acquired with a density-adapted 3D radial acquisition technique (24): TE= 0.4 ms, TR=11 ms, $T_{RO}$= 7.1 ms, flip angle= 36°, number of projections=10000, number of averages=10, voxel size=(6 x 6 x 6 mm)$^3$, total acquisition time=18:20 min.
For $^1$H and $^{23}$Na MRI of the heart synchronization of the data acquisition with the cardiac cycle was achieved using an acoustic cardiac gating device using an MR compatible stethoscope (easyACT, MRI.Tools GmbH, Berlin, Germany) (25-28).

**Data reconstruction and image analysis**

For 3D radial $^{23}$Na acquisitions the slice positioning given by the $^1$H images was used to extract the standard cardiac views from the reconstructed 3D dataset. Three reconstruction modi were applied for the 3D datasets obtained with the density-adapted 3D radial acquisition technique:

- ungated reconstruction of the acquired data
- retrospectively gated reconstruction using a temporal resolution of 0.1 s and time increments of 0.05 s. With this approach $^{23}$Na CINE images with 20 frames per cardiac cycle were reconstructed from the 3D acquisitions.
- retrospectively gated and cardiac phase selective reconstruction for data acquired during diastole. Based on the trigger information data acquired during systole were discarded.

Signal-to-noise ratio (SNR) maps were estimated using method 4 proposed in the NEMA standard MS-1 (29). According to (29) the noise in magnitude images is given by the standard deviation of the signal in a noise-only region divided by the factor 0.66. To eliminate the impact of potential artifacts of the radial acquisition scheme in any region of the image, the standard deviation of noise was derived from a separate noise scan. Contrast-to-noise-ratio (CNR) values were estimated by subtraction of the mean SNR value inside myocardium from the mean SNR of the left ventricular blood
pool. To examine the SNR profile along a circular trajectory inside the myocardium mid-ventricular short axis SNR maps were analyzed. For this purpose a standardized myocardial segmentation and nomenclature for tomographic imaging of the heart was used (30).
Results

**Hardware and RF characteristics**

The loaded ($Q_L$) and unloaded quality factor ($Q_U$) were obtained by loading and unloading the RF coil on a subject. The $^{23}$Na loop elements provided an average $Q_L/Q_U$ ratio of 0.12, indicating the dominance of sample noise. Reflection coefficients of -15 dB or less were observed. Element coupling was below -20 dB for the neighboring elements. Anterior posterior coupling was below noise level. The reflection coefficient of the $^1$H dipole was -26 dB. The decoupling of the $^{23}$Na loop elements to the $^1$H dipole was below -49 dB at 78.6 MHz and below -26 dB at 297 MHz for all $^{23}$Na loop elements. The complete set of S-parameters for both frequencies is surveyed in Figure 2 a, b.

Overall losses from the system’s RF output to the individual coil connectors were determined to be -0.7 dB for the $^{23}$Na RF chain and –1.4 dB for the $^1$H path, which were considered for the RF input power settings.

**EMF and SAR Simulations**

Figure 2c shows the combined transmit field for the $^{23}$Na array driven with the CP like phase setting for a transversal slice through the human voxel model’s heart. SAR$_{10g}$ values derived from the EMF simulations are outlined in Figure 2d. For the CP like mode the maximum local SAR$_{10g}$ per input power was found to be 0.39 1/kg. For the $^1$H dipole element the transmit field is shown in Figure 2e. The local SAR$_{10g}$ assessment yielded a local SAR$_{10g}$ per input power of 1.46 1/kg. With an input power of 25 $W_{RMS}$ for
the $^{23}$Na array and 6.8 $W_{RMS}$ input power for the $^1$H dipole and with considering the measured power losses in the RF chain, SAR$_{10g}$ values were below the limits permitted by the IEC guidelines (21). The first and second level controlled mode given by the IEC guidelines were not used for in vivo imaging which limits the maximum local SAR$_{10g}$ to 10 W/kg and preserves a safety margin of factor 2.

**Cardiac MR imaging**

The integrated dipole element supported basic $^1$H MR for localization as illustrated in Figure 3a. This enabled shimming planning of standard cardiac views in a clinical workflow for subsequent $^{23}$Na imaging. For $^{23}$Na MRI the transmitter reference amplitude was found to be 680±11 V, leading to a 180° pulse with a rectangular pulse applied for 1 ms.

For 3D gradient echo based $^{23}$Na MRI of the heart a nominal in-plane resolution of (5 x 5) mm$^2$ and a slice thickness of 15 mm was found to be feasible. Figure 3b depicts a short axis view of the heart derived from 3D gradient echo based imaging. The cardiac SNR evaluated over all volunteers was found to be 15±4 in the blood pool and 10±3 in the myocardium. A detailed SNR evaluation along the myocardial segments of a mid-ventricular short axis view is outlined in Figure 3c. A blood myocardium contrast of 4 was obtained. A high signal intensity caused by the high $^{23}$Na concentration of rib cartilage was observed (Figure 3b).

For whole heart coverage 3D density-adapted radial $^{23}$Na acquisitions a nominal spatial resolution of 6 mm isotropic was achieved. This improvement versus 3D conventional gradient echo acquisitions helped to reduce partial volume effects along
the slice direction and enabled retrospective image reconstruction of standard or arbitrary views of the heart. Figure 4a shows a whole heart coverage short axis view of the heart reconstructed from 3D radial acquisitions without using cardiac gating. A stack of short axis views covering the heart form the apex to the base shows consistent image quality for all slices reconstructed. Averaged over all subjects, mid-ventricular short axis views (Figure 4b) exhibited a SNR of 38±4 for the blood pool while myocardium revealed an SNR of 25±4. This provided a mean blood/myocardium contrast of 13. An exemplary SNR evaluation along the myocardial segments of a mid-ventricular short axis view shown in Figure 4b is illustrated in Figure 4c. In companion, a four chamber view of the heart reconstructed from the same 3D data set is shown in Figure 4d.

Figure 5 illustrates the sodium CINE imaging capabilities achieved with the proposed RF coil configuration in conjunction with 3D radial acquisitions and cardiac gating. Figure 5 a,b show end-systolic and end-diastolic short axis views of the heart derived from a CINE dataset consisting of 20 cardiac phases. To illustrate the movement and contrast throughout the cardiac cycle, a signal intensity profile along the long axis of the heart was extracted for every cardiac phase and summarized in the M-mode like view shown in Figure 5c.

Figure 5 d-e illustrates the impact of cardiac gating on the delineation of the myocardium. Figure 5d shows an un-triggered reconstruction. Figure 5e depicts the result of a cardiac gated reconstruction, based on data acquired during diastole only. The difference map in Figure 5f reveals a higher signal for the cardiac gated reconstruction inside the blood pool for regions in proximity to the myocardium. This
results in a blood-myocardium contrast enhancement of 20% versus the same data being reconstructed without cardiac gating.
Discussion

This feasibility study adds to the literature by demonstrating the feasibility of \(^{23}\text{Na}\) MRI of the human heart at 7.0 T. The evidence herein suggests that \(^{23}\text{Na}\) MRI at 7.0 T provides sensitivity and spatiotemporal resolution advantages over studies at 1.5 T and 3.0 T (10-12). The proposed RF coil setup exhibits adequate RF characteristics, transmission field homogeneity and penetration depth that support sodium imaging of the heart at 7.0 T in clinically acceptable scan times including free breathing, cardiac gated and ungated acquisitions. Depending on the acquisition technique used the SNR of myocardium was found to be between 10 and 27 for mid-ventricular short axis views of the heart. This achievement supports clear delineation of the myocardium including deep lying regions of the heart. The latter was reported to constitute a challenge for \(^{23}\text{Na}\) MR at a field strength of 3.0 T (31). Our in vivo studies in healthy subjects revealed that the use of a density-adapted 3D radial acquisition protocol used here provides an SNR advantage of roughly 80% compared to the used 3D gradient echo protocol. Pointing out this large gain we anticipate that differences in the calculation of the spatial resolution between Cartesian and radial acquisition protocols might slightly alter this value. The signal gain is related to the different echo times feasible for the 3D radial acquisition and the 3D gradient echo protocol due to the fast transverse relaxation of the sodium signal. Assuming a rapidly decaying component with \(T_{2,\text{rapid}}^*=(0.5-4.0)\) ms and a slowly decaying component with \(T_{2,\text{low}}^*=(12-20)\) ms (14), an SNR gain of 27% to 85% can be expected when TE is reduced from 1.9 ms to 0.4 ms. Additionally, different
pulse sequence parameters (e.g. TR) might contribute to the SNR gain. Yet, the sensitivity gain of the density-adapted 3D radial acquisition protocol enabled sodium CINE imaging of the heart with 10 frames per cardiac cycle within an acquisition time of approximately 19 min. Cardiac triggering can be considered being beneficial to enhance the delineation of myocardium and the blood pool for a diastolic cardiac phase, since the epi- and endocardial boarder sharpness was improved by not including data acquired during cardiac phases of myocardial contraction and relaxation. The 6 mm isotropic resolution of the whole heart 3D radial acquisitions facilitated the reconstruction of arbitrary cardiac views including a stack of short axis and four chamber standard cardiac views from a single acquisition.

Efficient and uniform RF excitation of the upper torso might present a challenge. This can be accomplished with a whole body RF coil as recently reported for $^{23}$Na MR at 3.0 T using a 16-rung birdcage resonator body coil (32). Since available transmit RF power is limited the high transmit power requirements of volume coils constrain the flip angles needed for short excitation pulses used in sodium MRI. The higher efficiency of a local transceiver array proposed in this work offsets this constraint since the applicable flip angles are restricted by SAR limitations rather than by RF peak power constraints. To our knowledge, this is the first cardiac $^{23}$Na MR study conducted with a multi-element transceiver coil.

The preliminary results reported for this feasibility study are likely to pave the way for further advances in RF coil technology tailored for assessment of myocardial sodium content. These efforts will help to further gain sensitivity by reducing loop element size and by including more loop elements and hence will contribute to further improvements
in spatio-temporal resolution. For example a two channel transmit loop together with a 8 channel receive array was reported to be feasible for sodium cardiac imaging at 3.0 T (33). The measured Q-ratio of 8 for the proposed four channel transceiver array leaves reasonable space for further subdivision of the elements into an array of eight or more TX/RX elements as recently demonstrated for transceiver arrays customized for $^1$H MRI at 7.0 T (34-41). To advance the proton imaging capabilities it stands to reason to incorporate multiple $^1$H elements into the anterior and posterior sections of the $^{23}$Na array to support high spatial resolution and accelerated imaging of the heart (42-48). This can either be achieved by dipole elements placed inside the loops or by overlapping loops for proton imaging as reported previously for a lower abdomen RF coil (49). For this purpose the bowtie radiative elements used here can be replaced by other dipole configurations including loopoles or bended dipoles (50,51).

We recognized limitations in our feasibility study, which included healthy volunteers before extra variances due to pathophysiological conditions are introduced. Taking into account that cardiac MR at 7.0 T is a field in a state of creative flux we felt that it is important to begin by reporting on the details of our preliminary results deduced from $^{23}$Na MRI of the heart before the technology will be placed in the hands of a broader group of clinical colleagues. Because of the physiological relevance of the intra-versus the extracellular sodium compartment, a separation between these pools constitutes another goal for the development of $^{23}$Na-MRI techniques employing relaxation-weighted imaging, multiple-quantum filtering (52-56) or bi-exponential-weighted imaging (57,58). Explorations into appropriate suppression or selective excitation techniques would be beneficial for suppression of signal contributions from
blood and cartilage which would help to enhance the dynamic range available for myocardial signal. For example, the cartilage signal could be spoiled by using a local surface gradient coil placed around the stenum on top of the chest. We are aware that spill-over effects from signal contributions of the blood pool might still constitute a challenge for myocardial sodium assessment in routine clinical applications of $^{23}$Na MRI. In analogy to recent pioneering reports on $^{23}$Na MR of the brain iterative reconstruction techniques that consider information from $^1$H imaging could be utilized to improve spatial resolution for sodium imaging of the heart at 7.0 T (59). Additionally, the blood signal could be eliminated by using inversion-recovery or relaxation-weighted imaging. For quantitative assessment of myocardial tissue sodium concentration, partial volume correction methods might be applied. For this purpose the geometric transfer matrix algorithm might be used (60). This algorithm was designed for positron emission tomography and recently transferred to non-proton MRI (61).

With the preliminary results achieved in this study we anticipate to make use of the proposed RF configuration for driving translational research. While being important but beyond the scope of this work, application of the proposed $^{23}$Na MRI setup could benefit the assessment of sodium homeostasis and myocardial viability (2,3,5). Furthermore, $^{23}$Na MRI at 7.0 T could provide means for the examination of cardiac sodium channelopathies (62). It is also clinically appealing to pursue sodium MRI of the heart at 7.0 T to gain better insights into hypertrophic cardiomyopathy (HCM). Previous animal studies reported a decrease in Na$^+$,K-ATPase activity for HCM, which resulted in a 40% increase in intracellular sodium concentration (63). $^{23}$Na MR at 7.0 T could be
also beneficial for scrutinizing alterations in the myocardial Na⁺ concentration evoked by cardiac drugs acting upon sodium ion channels (64-66).

Quantification of myocardial sodium concentration requires correction of coil-array sensitivity variation over the heart. This can be accomplished with B₁ mapping to derive the coil array’s transmission field distribution and the coil array’s sensitivity profile as demonstrated for various RF coil configurations tailored for ²³Na or other x-nuclei MRI at 3.0 T and 7.0 T (67-69). Improving in-plane spatial resolution of sodium MRI may be another important aspect to support these clinical applications. While the proposed RF coil design is tailored to the heart it can be adapted to support in vivo assessment of human sodium content in body sections other than the heart including the kidney or skin (67,70).

To conclude, the proposed transceiver array enables sodium imaging of the human heart at 7.0 T within clinical acceptable scan times and provides encouragement for further explorations into densely packed multichannel transceiver coil arrays tailored for ²³Na cardiac MR. The benefits of such improvements would be in positive alignment with the needs of explorations that are designed to examine the potential of ²³Na MRI for the assessment of myocardial ischemia and cell integrity. We suggest that ²³Na CMR at 7.0 T can also help to unlock questions regarding Na⁺ balance and Na⁺ storage functions of myocardial tissue with the ultimate goal to provide imaging means for diagnostics and for guiding treatment decisions in cardiovascular and metabolic diseases.
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References


Figure 1:
Overview of the proposed $^1$H/$^{23}$Na RF coil array tailored for cardiac $^{23}$Na MR at 7.0 T. **a)** Anterior and posterior coil section of the proposed sodium RF coil array placed on a mannequin. **b)** Basic loop structure and element numbering of the sodium array shown in the simulation model. **c)** Circuit diagram of the anterior section of the sodium loop array. **d)** Picture photograph of the dipole element used for proton imaging, which was placed inside loop element two as illustrated in b). **e)** Simulation setup consisting of the sodium array, the dipole element and the voxel model Duke inside the bore given by the gradient shield.
Figure 2:

Synopsis of the results derived from EMF simulations and measurements of the RF characteristics of the $^{23}$Na/$^1$H setup. **a**) Scattering parameters measured on a subject to demonstrate the resonance behavior of the $^{23}$Na/$^1$H setup at 78.6 MHz. **b**) Scattering parameters measured on a subject at 297 MHz. **c**) Transmission field simulation for the four channel $^{23}$Na array using a circular polarized phase setting depicted in a transversal plane through the heart marked by the ROI shown in red. **d**) Maximum projection plot of the local SAR$_{10g}/P_{\text{Input, RMS}}$ distribution of the $^{23}$Na array simulated for the human voxel model Duke based on the circular polarized phase setting. **e**) Transmission field simulation at 297 MHZ for the $^1$H dipole element depicted in a transversal plane through the heart (ROI in red). **f**) Maximum projection plots of the local SAR$_{10g}/P_{\text{Input, RMS}}$ distribution of the $^1$H dipole element simulated for the human voxel model Duke.
Figure 3:

a) $^1$H four chamber view localizer of the heart derived from 2D CINE gradient echo imaging using the dipole element. This systolic view was used for planning the 3D slabs employed in $^{23}$Na MRI, which were centered around a mid-ventricular short axis view marked by the orange line. b) $^{23}$Na short axis view of the heart acquired with the four channel loop array at the mid-ventricular slice position indicated in the $^1$H four chamber view shown in a). A 3D gradient echo imaging protocol was applied for data acquisitions. c) SNR evaluation for the mid-ventricular short axis view shown in b) following a trajectory along the myocardial segments 7-12. To guide the eye, the standardized myocardial segment model proposed in (30) is illustrated in orange.
Figure 4:
Cardiac $^{23}$Na images derived from ungated reconstruction of a 3D dataset which was acquired with a nominal isotropic spatial resolution of 6 mm using the 3D density-adapted radial acquisition protocol. Standard cardiac views were extracted using normal vectors from $^1$H image positioning. a) A stack of short axis views covering the whole heart from the apex to the atrium. b) Mid-ventricular short axis view used for SNR evaluation. c) SNR evaluation for the mid-ventricular short axis view shown in b) following a trajectory along the myocardial segments 7-12. d) Four chamber view of the heart retrospectively reconstructed from the 3D whole heart coverage data set.
Figure 5:
Cardiac CINE images reconstructed from a 3D dataset with a nominal isotropic spatial resolution of 6 mm which was acquired with the 3D density-adapted radial protocol. Retrospective gating was applied during reconstruction using time increments of 0.05 s and time frames of 0.1 s. Short axis views were extracted using the normal vector from $^1$H image positioning. 

a) Mid-ventricular short axis CINE frame in diastole. b) Mid-ventricular short axis CINE frame in systole. c) M-mode like, temporally resolved signal evolution extracted from the profile placed along the long axis of the heart, as depicted in a) and b), for all 20 frames of the CINE acquisition. The line shown in a) and b) are marked by the arrows A and B. d) Ungated reconstruction based on the complete dataset. e) Selective reconstruction of diastolic data acquired between 0.6 s and 1.2 s after recognition of the onset of the cardiac cycle using the acoustic cardiac triggering device. f) Difference map of images deduced from ungated and gated reconstruction which revealed a blood pool signal gain of +20% of the gated image. This behavior helped to enhance the delineation between blood and myocardium.