Normal Values of Magnetic Relaxation Parameters of Spine Components with the Synthetic MRI Sequence

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ABSTRACT

BACKGROUND AND PURPOSE: SyMRI is a technique developed to perform quantitative MR imaging. Our aim was to analyze its potential use for measuring relaxation times of normal components of the spine and to compare them with values found in the literature using relaxometry and other techniques.

MATERIALS AND METHODS: Thirty-two spine MR imaging studies (10 cervical, 5 dorsal, 17 lumbosacral) were included. A modified multiple-dynamic multiple-echo sequence was added and processed to obtain quantitative T1 (millisecond), T2 (millisecond), and proton density (percentage units [pu]) maps for each patient. An ROI was placed on representative areas for CSF, spinal cord, intervertebral discs, and vertebral bodies, to measure their relaxation.

RESULTS: Relaxation time means are reported for CSF (T1 = 4273.4 ms; T2 = 1577.6 ms; proton density = 107.5 pu), spinal cord (T1 = 780.2 ms; T2 = 101.6 ms; proton density = 58.7 pu), normal disc (T1 = 1164.9 ms; T2 = 101.9 ms; proton density = 78.9 pu), intermediately hydrated disc (T1 = 723 ms; T2 = 66.8 ms; proton density = 60.8 pu), desiccated disc (T1 = 554.4 ms; T2 = 55.6 ms; proton density = 47.6 pu), and vertebral body (T1 = 515.3 ms; T2 = 100.8 ms; proton density = 91.1 pu). Comparisons among the mean T1, T2, and proton density values showed significant differences between different spinal levels (cervical, dorsal, lumbar, and sacral) for CSF (proton density), spinal cord (T2 and proton density), normal disc (T1, T2, and proton density), and vertebral bodies (T1 and proton density). Significant differences were found among mean T1, T2, and proton density values of normal, intermediately hydrated, and desiccated discs.

CONCLUSIONS: Measurements can be easily obtained on SyMRI and correlated with previously published values obtained using conventional relaxometry techniques.

ABBREVIATIONS: PD = proton density; pu = percentage units

MR imaging provides excellent soft-tissue contrast. However, this information is qualitative; on T1WI or T2WI, the absolute intensity has no physical unit and diagnosis relies on relative contrast differences between adjacent tissues in the image.1 Nevertheless, quantitative imaging is rapidly gaining importance as a complement to qualitative sequences, such as diffusion-based sequences, perfusion imaging, and MR spectroscopy.

Relaxometry is a quantitative method that allows the absolute quantification of MR imaging parameters such as T1 and T2 relaxation times as well as proton density (PD). This technique is nevertheless very time-consuming and requires T1 and T2 quantifications to be performed separately through different types of sequences that were not feasible from a practical point of view in a clinical routine setup.2 These sequences included inversion recovery3 for T1 measurements or techniques derived from modifications to the Carr-Purcell Meiboom-Gill sequence, considered the criterion standard for the measurement of T2 relaxation time.4,5 Several methods are currently being developed to perform quantitative MR imaging techniques in a more efficient manner. MR fingerprinting consists of a pseudorandomized variation of the sequence parameters (flip angle, TR, and k-space trajectory), thus producing a unique signal evolution or “fingerprint” for each tissue.6 Following acquisition, the process involves a pattern-
identification algorithm to link the obtained fingerprints to those stored in a predefined dictionary. Recent studies aim to differentiate common types of adult intra-axial brain tumors or to perform quantitative abdominal imaging using this method.

Another MR imaging quantitative method is the SyntAc sequence (based on the QRAPMASTER sequence, SyntheticMR, Linköping, Sweden). This is the technique we have chosen for our study because it is more readily available and is already approved by the Conformité Européenne (CE) and FDA. In this method, physical properties (PD, longitudinal [T1] and transverse [T2] relaxation rates, and correction for B1-inhomogeneities) are calculated using a single, multiecho, saturation recovery TSE sequence. Subsequently, any “synthetic” T1WI or T2WI can be achieved from the absolute parameters.

Common advantages of these 2 methods for brain imaging are reduced acquisition time and the possibility of performing automatic tissue segmentation and volume estimation. This has already been used for the evaluation of demyelinating diseases. The recent study published by Tanenbaum et al compared synthetic-versus-conventional MR imaging for routine neuroimaging, concluding that synthetic MR imaging quality was comparable with conventional sequences despite artifacts being more common in the synthetic FLAIR sequence, as previously reported. This shortcoming was also reported with MR finger imaging, concluding that synthetic MR imaging quality was comparable with conventional sequences despite artifacts being more common in the synthetic FLAIR sequence, as previously reported.

The goal of this study was to measure T1 and T2 relaxation times, as well as PD values, of normal structures included in normal spine MR imaging (CSF, spinal cord, healthy-versus-desic-cated intervertebral disc, and vertebral body) using Synthetic MRI (SyMRI) 8.0 software (SyntheticMR) (this method has already been used for quantifying relaxation times in the brain, but, as far as we know, it has not been used in the spinal region). Furthermore, we aimed to compare our values with those found in the literature from studies using relaxometry.

**MATERIALS AND METHODS**

**Participants**

Thirty-two nonconsecutive patients (17 males; 15 females; mean age, 55.47 years; range, 17–85 years) undergoing spine MR imaging studies (10 cervical, 5 dorsal, 17 lumbaroscal) at our institution (Geneva University Hospitals) in 2016 were included in this study.

All subjects met the following criteria: 1) a synthetic MR image added to our standard imaging protocol when time constraints allowed it, and 2) no history of neoplastic conditions, radiation therapy, or multiple sclerosis.

The study was approved by our institutional review board, and informed consent was waived.

**Image Acquisition**

All studies were performed on a 1.5T Ingenia scanner (Philips Healthcare) using a 52-channel spine coil. The imaging protocol included quantitative sagittal MR imaging and the conventional MR imaging sequences used in our center, which include at least conventional 2D sagittal spin-echo T1-weighted, FSE T2-weighted, and STIR sequences.

A multiple-dynamic multiple-echo sequence (SyntAc) was modified for spine imaging. Parameters were the following: sagittal orientation, FOV = 200 × 321 mm, acquisition (reconstruction) voxel size = 0.89 × 1.48 mm² (0.71 × 0.72 mm²), 15 contiguous slices of 4-mm thickness, TE = 11/100 ms, TR = 2485 ms, TSE factor = 12, sensitivity encoding acceleration factor = 2. The acquisition time of the synthetic sequence was 5 minutes 40 seconds.

This sequence generates 8 different contrasts (2 different TEs and 4 different TIs), which are then processed by SyMRI 8.0 software to obtain quantitative T1 (millisecond), T2 (millisecond), and PD (pu) maps. The PD maps are measured in percentage units (pu), in which 100% is the value of brain CSF that is used for calibration. Because fat has a higher signal intensity than CSF, it is displayed with a PD value above 100 pu (0–108 pu if T1 > 1100 ms and 0–120 pu if T1 < 1100 ms). The dynamic ranges of T1 and T2 values are 300–4300 and 20–2000 ms, respectively (this information was extracted from the software manual).

These maps were then saved in DICOM format and analyzed in OsiriX Imaging Software (Version 7.0.3; http://www.osirix-viewer.com).

**Radiologic Assessment**

All images were analyzed by a neuroradiologist and a fellow in neuroradiology on standard radiologic workstations connected to the PACS. A consensus was reached in the choice of representative normal-intensity areas for CSF, spinal cord, intervertebral disc, and vertebral body on each study (based on the conventional MR imaging sequences). We subsequently placed an ROI on these areas and measured representative T1 (millisecond), T2 (millisecond), and PD (pu) of each spinal element, carefully avoiding partial volumes (Figs 1 and 2).

The cervical cord was assessed at the C2 and C7 levels. The dorsal spinal cord was assessed on an upper and lower level around T2 and T10, respectively. In the lumbar spine, a single ROI was placed on the conus medullaris. We placed the ROI in the center of the spinal cord, including both gray and white matter, to elude a possible partial volume effect.

Two ROIs were placed in each spinal region for CSF, also on the highest and lowest possible positions, carefully avoiding descending nerve roots and flow artifacts. In the cervical studies, the upper ROI was placed in the cisterna magna, and the lower one, at the level of C7. In the dorsal studies, ROIs were placed at levels similar to the ones used for the spinal cord. In the lumbar studies, one ROI was placed at the level of the conus medullaris, and another, around S1.

All cervical, dorsal, and lumbar vertebrae fully within the FOV were included, as well as the first sacral vertebra. ROIs were placed on the anterior part of the vertebral body to avoid vascular artifacts, cortical bone, partial volume effects, and areas with signal abnormalities (eg, hemangiomas) on conventional MR imaging sequences. We excluded 3 fractured vertebrae and 2 with metallic artifacts.

All cervical, dorsal, and lumbar discs that were completely visualized were evaluated, excluding 1 with artifacts secondary to instrumentation. Intervertebral discs were measured and assigned a status of clearly normally hydrated (equivalent to a grade I of the Pfirrmann scoring), intermediate signal (grades II or...
III), or desiccated (grades IV or V) by visual assessment. The intermediate signal status included discs that were neither clearly normally hydrated nor clearly desiccated. Discs showing complete loss of height were not included.

These ROIs were originally placed on the sagittal T2-weighted image and copied and pasted into the sagittal synthetic T1, T2, and PD maps; their mean value, SD, and area were calculated.

**Validity of Quantification Measurements**

To verify whether the quantification with our modified SyMRI sequence with a relatively low TR (2485 ms) was correct, we compared the T1, T2, and PD values obtained on a set of gel tubes composed of different agar and gadolinium concentrations measured with the SyMRI sequence with a TR of 6000 ms (and we used this technique as the criterion standard). Apart from TR, all parameters were kept identical. The acquisition time of this sequence was 11 minutes 20 seconds, which precludes its use in clinical routine. We also added an idealized model with mineral water to the FOV to include a compartment with high T1 and T2 values.

**Statistical Analysis**

Means and standard errors of the mean of T1, T2, and PD values for CSF, spinal cord, discs (normal, desiccated, and intermediate hydration), and vertebral bodies across all spinal levels were calculated. A 1-sample Kolmogorov-Smirnov test was used to investigate whether continuous variables were normally distributed across groups (spinal region or disc hydration status). If normal distribution of the data was confirmed in all groups, differences between group mean values were investigated by 1-way ANOVA with age as a covariate using a model with type III sum of squares. Pair-wise comparisons were subsequently performed by the least significant difference method. If data were not normally distributed in at least 1 group, differences among group median values were investigated by the nonparametric Kruskal-Wallis H test. The Mann-Whitney U test was used for pair-wise comparisons in this case.

Correlation coefficients ($R$) were calculated, and linear regression was performed between the values obtained with the SyMRI sequence used in the study (TR = 2485 ms) and those obtained by the criterion standard (TR = 6000 ms).

A .05 two-tailed significance level was used for all tests. Statistical analysis was performed with the Statistical Package for the Social Sciences, Version 22 (IBM, Armonk, New York).
RESULTS

Table 1 shows the linear regression and the correlation coefficients obtained for T1, T2, and PD values measured with our SyMRI sequence and with a sequence with a long TR (6000 ms) used as the criterion standard. Correlation was high for T1 and T2 values ($R = 1.00$ for both) and moderately high for PD values ($R = 0.57$). The slope of the line of best fit was close to 1 in all 3 analyses (1.06, 0.96, and 0.94 for T1, T2, and PD, respectively).

Descriptive statistics with means and 95% confidence intervals of the mean of T1, T2, and PD values of CSF, spinal cord, normal discs, intermediately hydrated discs, desiccated discs, and vertebral bodies for all spinal levels are shown in Table 2.

Comparison of Spinal Levels

The Kolmogorov-Smirnov test showed that only T1 and PD values for CSF were not normally distributed in any of the spinal levels (cervical, dorsal, lumbar, and sacral) ($P < .05$). The non-parametric Kruskal-Wallis H test was therefore used to compare medians of spinal levels in these 2 variables. The distribution of all the other variables was not significantly different from normality in all spinal levels, which allowed comparison of means by ANOVA.

We did not find a significant difference among different spinal levels in spinal cord T1 values ($P = .436$). The T1 value of normal discs across spinal levels was significantly different, with cervical spine discs showing the largest difference in relation to dorsal and lumbar discs ($P < .05$). There was an even more significant effect of spinal level on the T1 value of vertebral bodies ($P < .001$). These results are summarized in Fig 3.

Spinal level had a significant effect on T2 signal intensity of the spinal cord and normal discs ($P < .05$ and $P < .001$, respectively). Overall, the T2 value of normal discs increased progressively from the cervical to the lumbar region. The T2 value of vertebral bodies was not significantly affected by spinal level ($P = .370$) (Fig 4). Similarly, CSF T2 values did not show a significant difference among spinal levels ($P = .081$) (Fig 5).

PD values of spinal cord, normal discs, and vertebral bodies were significantly affected by spinal level ($P < .01$, $P < .001$, and $P < .001$, respectively). PD values of the spinal cord and vertebral bodies tended to decrease from cervical to lumbar, whereas PD values of discs tended to increase (Fig 6).

Medians of CSF T1 values were not significantly different across spinal levels ($P = .234$). Despite the noticeable similarity of CSF PD values in the 4 different spinal levels, a significant difference was found among medians ($P < .05$) (Table 3).

Comparison of Disc Hydration Statuses

Means of disc T1 values were strongly affected by disc hydration status ($P < .001$), with progressively lower values as the disc water content decreased (Fig 7). The same trend was found on the non-parametric test results with disc T2 and PD values significantly decreasing as disc hydration declined ($P < .001$) (Table 4).

DISCUSSION

CSF

Our results for CSF mean relaxation times are comparable with those found in the literature. Krauss et al19 showed T1, T2, and PD values very similar to ours on the charts displaying their phantom measurements with the QRAPMASTER technique. Using the same technique, Warnijes et al1 obtained T1 and T2 values of 3940 ms and 1910 ms, respectively, and PD values of 1029 pu, specifying that “the PD of pure water at 37°C corresponds to 1000” and arguing that the proton density of CSF appeared “somewhat high, possibly due to flow-effects or diffusion.” In a review article from 2017, Bojorquez et al14 reported the following CSF T1 values in
There is also variation in the total number of tracts (which differ from our reported T2 values for CSF across spinal levels assuming a fixed age of 55.5 years, respectively (single asterisk indicates \( P < .05 \); double asterisks, \( P < .01 \); triple asterisks, \( P < .001 \)). Significant pair-wise differences between spinal levels are indicated by a different letter assigned to each level.

![FIG 4. ANOVA results with comparison of estimated marginal means of T2 values for spinal cord, normal discs, and vertebral bodies across spinal levels, assuming fixed ages of 56.2, 37.2, and 56.4 years, respectively (single asterisk indicates \( P < .05 \); double asterisks, \( P < .01 \); triple asterisks, \( P < .001 \)). Significant pair-wise differences between spinal levels are indicated by a different letter assigned to each level.](image)

The T2 relaxation times reported in the literature are 73 ms for white matter and 76 ms for gray matter, which differ from our T2 mean value of 101.6 \( \pm \) 2.9 ms. Lu et al. reported T2 values for brain gray matter at 1.5T, ranging from 81 (putamen) to 99 ms (cortex). We did not find any previously reported PD values for the spinal cord.

Although the absolute values did not show large variations, we found that the cervical region had a significantly different T2 value from the thoracic and lumbar regions (94.1 versus 104.4 and 103.6 ms, respectively). The same difference was found in PD values (63.3 versus 58.7 and 55.7 \( \mu \)u, respectively). No significant differences were found for T1 values. This may relate to the normal anatomy of the spinal cord because the configuration of the gray matter and gray/white matter ratio varies along spinal cord regions, depending on the neural requirements of a given region. There is also variation in the total number of tracts (which decreases from cervical to lumbar, as more and more nerve projections arise), as well as in the cross-sectional area of the cord (which increases at the cervical and lumbar enlargements). These 2 features determine variations in the density of spinal tracts along different segments of the cord.

We believe SyMRI could prove to be a good quantitative MR imaging technique for the assessment of the spinal cord. T1, T2, and PD measurements have the potential to be used as biomarkers of both myelin and axonal integrity. A promising application could therefore be to evaluate the degree of spinal cord integrity in posttraumatic or demyelinating lesions. Further studies would be needed to corroborate the variations we found among spinal levels and clarify the possible causes. Recently, the role of T1, T2, PD, and myelin measurements has been analyzed in different neurologic conditions in the brain, and we believe it could potentially be directly translated into spinal cord conditions. Moreover, although outside of the scope of this study, myelin water fraction can be extracted from SyMRI T2 measures. Preliminary studies...
Discs
We have found significant differences among mean T1, T2, and PD values of normal, intermediate, hydrated, and desiccated discs, confirming the formerly described negative relationship between relaxation time (T1, T2) and disc-degenerative grade. T1 relaxation time has been shown to decrease with disc degeneration and herniation.

T2 relaxation times are related to disc biochemical composition in that decreased T2 values represent decreased disc water content and proteoglycan loss or replacement of the nucleus by degenerative tissue, which has lower T2 values.

Our results also showed significant differences in disc values among different spinal levels, with lower T1, T2, and PD values in the cervical spine compared with the thoracic or lumbar levels. This finding agrees with previous biochemistry reports of lower water content in cervical discs compared with lumbar discs.

Another possible explanation for our significantly lower values in cervical discs could be related to the ROI areas, which were smaller in that region. This feature may have affected the proportion of anulus fibrosus or its analogous fibrous structure, the intranuclear cleft versus the nucleus pulposus. Because the reduction in cervical disc areas is mainly at the expense of nucleus pulposus, this finding could be another reason for their decreased T1, T2, and PD values.

Limitations to our conclusions in disc analysis could be because we established disc hydration status by agreed visual assessment and the age range is wide in our population (17 to 81 years), possibly explaining the large variability of the values obtained for discs (and vertebral bodies).

Disc degeneration is a condition reported daily on spine examinations, and SyMRI could help establish universal cutoffs (settled by spinal level) for staging it.

Vertebral Bodies
Our mean values of T1 and T2 mean relaxation times are consistent with those found in the literature. In the Bojorquez et al review, the values obtained for T1 vary between 106 ± 4 ms and

### Table 3: Results of the Kruskal-Wallis H test with T1 and PD median values for CSF across spinal levels

|                | Cervical | Dorsal | Lumbar | Sacral | P   |
|----------------|----------|--------|--------|--------|-----|
| T1 CSF         | 4287.7   | 4285.5 | 4289.6 | 4294.3 | .234|
| PD CSF         | 108.0ab  | 108.0a | 108.0b | 108.0ab| <.05|

Note: Significant pairwise differences between spinal levels are indicated by a different letter (a, b) assigned to each level.

### Table 4: Results of the Kruskal-Wallis H test with median T2 and PD values for discs across the 3 hydration status

|                | Normal | Intermediate | Desiccated | P   |
|----------------|--------|--------------|------------|-----|
| T2 disc        | 95.5a  | 66.2b        | 54.0c      | <.001|
| PD disc        | 80.9a  | 58.0b        | 47.8c      | <.001|

Note: Significant pairwise differences between hydration status are indicated by a different letter (a, b, c) assigned to each level.
586 ± 73 ms,3 and from 40 to 160 ms for T2.16 Le Ster et al17 reported T1 values of 621 ms.

T2 relaxation times did not show significant differences among spinal levels, but T1 values significantly decreased from cervical to lumbar (from 547.1 to 495.1 ms) as did PD values (from 92.3 to 82.4 pu). Because bone marrow is characterized by large variations in T1 relaxation times between water and fat compartments,38 the differences found may be due to normal anatomic changes in the proportion of fatty marrow along the spine (which increases from C3 to L5).19

A potential application of determining T1, T2, and PD values of vertebral bodies could be in the detection of bone marrow infiltrative conditions, such as hematologic malignancies, by providing a quantitative tool for imaging assessment in situations in which it is difficult to visually confirm and demarcate pathologic changes. However, consideration must be given to the high variability relating to vertebral body composition among healthy individuals because it depends on different factors such as age, with conversion from red to yellow marrow, or the effect of hematocrit levels.

The On-line Table outlines previously published measured values with conventional relaxometry techniques for CSF, spinal cord, disc, and bone marrow.

Our study has some limitations, some of which have already been mentioned above. Another limitation is the potential bias introduced by manually performing some operations such as ROI selection. For spinal cord measurements, atlas-based techniques are available that could potentially have allowed gray-white matter distinction.

Nonconsecutive selection of patients may have introduced some bias. We nevertheless believe that this is minimized because there were no criteria in selecting patients for the study other than time constraints allowing the synthetic sequence to be performed. Taking this into account, all patients in whom this sequence was performed, within the time considered, took part in the study; this feature made the selection consecutive in this subgroup of patients.

We probably also have some limitations in accurately quantifying elements with high T1 and PD because our sequence has a relatively low TR (2485 ms). This limitation means that complete relaxation of high T1 structures is not fulfilled during the sequence; therefore, signal can be somehow saturated. Our attempt to verify the validity of quantification due to our short TR showed a high correlation of values for a T1 range of 300–822 ms as well as in the idealized model with T1 of 2819 ms. However, we did not compare the validity of the measurements with higher T1 as is the case with CSF; this difference may have biased the results obtained in this compartment due to the low TR.

Moreover, when we analyzed structures with small dimensions, such as the spinal cord, our quantitative values could have partial volume effects due to our slice thickness of 4 mm. This could be an explanation for the higher T2 values found in the spinal cord compared with previously published values in the literature.

Another limitation of our study is the lack of comparison among the values obtained with an internal criterion standard. These values would be more meaningful if compared with quantitative measures using conventional techniques in the same patients. Unfortunately, it is difficult in practice to add conventional sequences to the imaging protocol due to the additional acquisition time needed. Similarly, conventional T1 and T2 quantitative mapping techniques were not used for the agar phantom experiments. We are aware that SyMRI with a TR of 6000 is not a true criterion standard. However, quantification with SyMRI (TR ranging from 2900 to 5000 ms) has been validated in several previous publications.1,11,19,40–42

CONCLUSIONS

T1, T2, and PD measurements can be easily and rapidly obtained with SyMRI, a tool that could be readily available as a quantitative method in the spinal region, retrieving results comparable with those made previously by other quantitative techniques.

It is likely that the use of quantitative MR imaging will be implemented as part of the radiologist’s every day work. Setting the reference range for relaxation times of CSF, spinal cord, vertebral bodies, and discs in healthy individuals will therefore have a straightforward application. These measurements could be used in our future daily practice as a quantitative extra aid in the radiologic assessment of spinal conditions, such as infiltration of bone marrow in hematologic disease, demyelinating disease, and spondylodiscitis. They could also play a role in the classification of disc degeneration and the choice of treatment.

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