Responses of the Human Brain to Mild Dehydration and Rehydration Explored In Vivo by $^1$H-MR Imaging and Spectroscopy

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*AJNR Am J Neuroradiol* published online 17 September 2015

http://www.ajnr.org/content/early/2015/09/17/ajnr.A4508
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ABSTRACT

BACKGROUND AND PURPOSE: As yet, there are no in vivo data on tissue water changes and associated morphometric changes involved in the osmo-adaptation of normal brains. Our aim was to evaluate osmoadaptive responses of the healthy human brain to osmotic challenges of de- and rehydration by serial measurements of brain volume, tissue fluid, and metabolites.

MATERIALS AND METHODS: Serial T1-weighted and $^1$H-MR spectroscopy data were acquired in 15 healthy individuals at normohydration, on 12 hours of dehydration, and during 1 hour of oral rehydration. Osmotic challenges were monitored by serum measures, including osmolality and hematocrit. MR imaging data were analyzed by using FreeSurfer and LCModel.

RESULTS: On dehydration, serum osmolality increased by 0.67% and brain tissue fluid decreased by 1.63%, on average. MR imaging morphometry demonstrated corresponding decreases of cortical thickness and volumes of the whole brain, cortex, white matter, and hypothalamus/thalamus. These changes reversed during rehydration. Continuous fluid ingestion of 1 L of water for 1 hour within the scanner lowered serum osmolality by 0.96% and increased brain tissue fluid by 0.43%, on average. Concomitantly, cortical thickness and volumes of the whole brain, cortex, white matter, and hypothalamus/thalamus increased. Changes in brain tissue fluid were related to volume changes of the whole brain, the white matter, and hypothalamus/thalamus. Only volume changes of the hypothalamus/thalamus significantly correlated with serum osmolality.

CONCLUSIONS: This is the first study simultaneously evaluating changes in brain tissue fluid, metabolites, volume, and cortical thickness. Our results reflect cellular volume regulatory mechanisms at a macroscopic level and emphasize that it is essential to control for hydration levels in studies on brain morphometry and metabolism in order to avoid confounding the findings.

ABBREVIATIONS: HCT = hematocrit; OSM$_{\text{serum}}$ = serum osmolality

The regulation of body fluid balance inherently determines serum osmolality. Due to the high permeability of the cell membranes for water, the direction of water movement is determined by the osmotic gradient across the cell membrane between the intra- and extracellular space. Thus, serum hyperosmolality causes cell shrinkage, whereas hypo-osmolality induces cell swelling.

In response, the cell initiates regulatory changes to the opposite direction. Initially, cells quickly re-adjust their volume by transmembranous ion movements. Volume regulation by electrolyte shifts is limited because alterations of ion gradients across the cell membrane interfere with the structure and function of intracellular macromolecules and membrane transporters. To avoid the adverse effects of changes in ion composition, the cell uses organic osmolytes, instead of ions, which allow volume adjustment without compromising cell function. Organic osmolytes encompass polyalcohols like myo-inositol; methylamines like glycerylphosphocholine; amino acids; and derivatives like glutamine, glutamate, N-acetylaspartate and N-acetyl aspartylglutamate, and creatine and taurine.

Metabolic responses to fluid imbalance have primarily been...
studied in animals and patients with pathologies affecting serum osmolality. Previous brain morphometry findings in healthy subjects undergoing de- and rehydration have remained somewhat inconclusive and inconsistent in terms of the distribution of the changes and the structures affected. Kempston et al,7,10 Dickson et al,11 and Watson et al12 induced dehydration by thermal exercises. They reported no significant effect of dehydration on brain volume. Results on ventricular volume ranged from decrease12 to increase,7,10 and Dickson et al found no changes.11 Duning et al13 showed a 0.55% brain volume reduction after 16 hours of thirsting and a 0.72% increase after subsequent rehydration. Similarly, Streitbürger et al14 demonstrated an increase in brain volume on ingestion of 3–4 L of water; however, they failed to show an effect of dehydration on brain volume by 2 days of restricted water intake. In a study on ultramarathon runners, Freund et al15 described a reversible 6% cortical volume reduction with 2 months of daily running. Currently, there are no in vivo data on tissue water changes and associated morphometric changes involved in the osmoadaptation of normal brains. In this study, we examined healthy individuals during normo-, de-, and rehydration by MR spectroscopy and morphometry and analyzed the resulting changes of global brain volume, gray/white matter volume, cortical thickness, tissue fluid, organic osmolytes, and serum parameters.

MATERIALS AND METHODS

Ethics Statement

The study was approved by the local Medical Ethics Committee (Faculty of Clinical Medicine, University of Würzburg), and all participants gave written informed consent before enrollment. The procedures that followed were in accordance with the Declaration of Helsinki.

Participants

We studied 15 healthy volunteers (6 women, 9 men) who received an allowance for participating in the study. Female and male participants exhibited a narrow age range (mean, 27.67 ± 2.26 years) and body weight (mean, 74.53 ± 12.01 kg), which did not differ across sexes (P > .05, Wilcoxon rank sum test). All subjects were nonsmokers and free of medication. Exclusion criteria were current or past substance abuse other than nicotine and any other psychiatric, neurologic, and medical illness.

MR Imaging

All measurements were performed on a 3T Tim Trio MR imaging system (Siemens, Erlangen, Germany) by using a transmit/receive head coil.

TI-Weighted 3D Imaging and Quantitative TI-Mapping. For volumetric and surface-based morphometry, an isotropic (1.0 × 1.0 × 1.0 mm³) whole-brain T1-weighted MPRAGE sequence optimized for subsequent image data analysis was acquired (TR = 2300 ms; TE = 2.98 ms; magnetization preparation by nonselective inversion recovery TI = 900 ms; along with fat suppression/fast water excitation to reduce signal from fatty bone marrow, scalp, and dura; excitation flip angle = 9°; 160 sagittal slices; matrix size = 256 × 256).

For quantitative T1-relaxation-time mapping, a partial-brain FLASH (GE Healthcare, Milwaukee, Wisconsin) sequence (TR = 330 ms, TE = 1.8 ms, 22 axial slices, matrix size = 384 × 384, 0.5 × 0.5 × 3.0 mm³) was recorded at 2 flip angles (8° and 37°).

1H-MR Spectroscopy. Localized 1H-MR spectroscopy data were recorded by single-voxel spectroscopy measurements using a point-resolved spectroscopy sequence (TE = 30 ms, TR = 2000 ms). Automatic dynamic frequency correction was used for MR spectroscopy with least squares optimization across the main peaks of the water-suppressed spectra. For all subjects and each of the longitudinal point-resolved spectroscopy sequence measurements (compare “Protocol of Longitudinal Measurements”), a voxel volume of 20 × 20 × 20 mm³ placed in the left paracentral lobule was measured. As described earlier,16-18 placing the voxel with its inferior edge at the callosomarginal sulcus and with its posterior edge at the central sulcus allowed highly reproducible positioning. Because the voxel was placed in white matter mainly (Fig 1) systematic differences between the partial volume contribution of white matter, gray matter, and CSF were excluded (ANOVA, P > .23). No partial volume correction was performed. In vivo resonance changes were quantified by ensuring fixed scanner calibration by using in vitro references and by correcting for differences in coil load. Spectra with and without chemical shift selective water suppression were obtained (80 versus 16 acquisitions) to allow eddy current correction and to quantify changes of brain tissue water in the point-resolved spectroscopy sequence voxel by using the spectra without water suppression.19,20

Protocol of Longitudinal Measurements

The study consisted of 2 MR imaging examinations measuring healthy volunteers on normo-, de-, and rehydration. The first examination studied the regularly hydrated participants, yielding a baseline 1H-MR spectroscopy and 3D T1-weighted dataset (normohydration). Normohydration at the time of the first examination was confirmed by blood and serum parameters (Table). Then, subjects were asked not to drink, not to eat meals containing more than 0.5 L of fluid, and to refrain from physical activity for 12 hours (dehydration). Subsequently, MR imaging measurements of the second examination included a 1H-MR spectroscopy and 3D T1-weighted dataset with subclinically dehydrated subjects. Participants then remained within the MR imaging scanner without being repositioned, and water was orally ingested via a plastic tube for 60 minutes at a rate of 1 L per hour (by using a MRidium MR imaging Infusion System; IRadimed, Winter Park, Florida) (dehydration). Simultaneously, 19 serial 1H-MR spectroscopy datasets and 1 final 3D T1-weighted structural scan were recorded during 67 minutes 4 seconds ± 6 minutes 45 seconds. To control for circadian hormone effects, we scheduled the first MR image in the evening and the second in the morning of the following day for every participant.

For measuring serum parameters, 3 blood samples were drawn consecutively at normo-, de-, and rehydration via an antecubital IV line. Mean corpuscular hemoglobin concentration and hematocrit (HCT) values were determined by an automated counter using the electrical impedance method (Counter S-Plus IV; Beckmann Coulter, Fullerton, California).

Serum electrolytes were measured by a potentiometric autoanalyzer (Ektachem 700XRC; Eastman Kodak, Rochester, New York).
**Data Postprocessing**

MR Morphometry. Morphometric analyses were performed to detect changes of whole brain, white matter, subcortical and cortical gray matter volume, and cortical thickness. To extract reliable cortical volume and thickness estimates, we automatically processed images by using the longitudinal stream within FreeSurfer (Version 5.3.0; http://surfer.nmr.mgh.harvard.edu). For details, please see the Online Appendix.

**Blood/serum parameters during normo-, de-, and rehydration**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Normohyd</th>
<th>Dehyd</th>
<th>Rehyd</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na⁺ (mmol/L)</td>
<td>137.1 ± 2.7</td>
<td>138.5 ± 2.2</td>
<td>136.9 ± 2.3</td>
</tr>
<tr>
<td>K⁺ (mmol/L)</td>
<td>4.6 ± 0.4*</td>
<td>5.2 ± 0.6*</td>
<td>4.9 ± 0.4*</td>
</tr>
<tr>
<td>HCT (%)</td>
<td>41.9 ± 2.9</td>
<td>43.8 ± 2.9</td>
<td>42.5 ± 2.9</td>
</tr>
<tr>
<td>MCHC (g/dL)</td>
<td>34.3 ± 0.9</td>
<td>34.3 ± 0.8</td>
<td>34.3 ± 0.8</td>
</tr>
<tr>
<td>Urea (mg/dL)</td>
<td>27.4 ± 6.2</td>
<td>26.8 ± 5.9</td>
<td>25.5 ± 5.6</td>
</tr>
<tr>
<td>OSMserum (mOsm/kg)</td>
<td>312.7 ± 4.8*</td>
<td>314.8 ± 4.1*</td>
<td>311.8 ± 4.7*</td>
</tr>
<tr>
<td>Glucose (mg/dL)</td>
<td>92.1 ± 6.8*</td>
<td>84.4 ± 6.9*</td>
<td>85.5 ± 5.9*</td>
</tr>
</tbody>
</table>

**Note:**—Normohyd indicates normohydration; Dehyd, dehydration; Rehyd, rehydration; MCHC, mean corpuscular hemoglobin concentration.

* Significant changes among different levels of hydration (normo-, de-, and rehydration) at type I error probabilities of P < .05 (1-sided t tests; df = 14).

The integral of unsuppressed water was used as a measure of brain tissue water (H₂O_brain). Data of each metabolite and of brain tissue fluid were normalized (by subtraction) to the second session (dehydration) by using custom-written Matlab scripts (MathWorks, Natick, Massachusetts). Outliers were defined as values exceeding 2.5 SDs, and data were corrected for outliers. Units were expressed as normalized percentage signal change. Thus, dehydration was chosen as the reference to conveniently plot successive changes (compare “MR Morphometry” above).

Criteria for assessment of spectral quality were the following: 1) the signal-to-noise ratio (defined by the ratio of the maximum in the spectrum-minus-baseline over the analysis window to twice the root-mean-square residuals, 2) the line width (roughly estimated by using the full width at half maximum), 3) the distribution of residuals, and 4) the SD. In the LCMedal, SD estimates are Cramer-Rao lower bounds. Cramer-Rao lower bounds of >50% indicate that the metabolite concentration may range from zero to twice the estimated concentration. In this study, we selected for further analysis only spectral data meeting the following: 1) a signal-to-noise ratio of >4; 2) full width at half maximum of ≤0.07 ppm; 3) randomly distributed residuals of ~0; and 4) Cramer-Rao lower bounds of <25%.

**1H-MR Spectroscopy.** 1H-MR spectroscopy data were analyzed by using the frequency-domain-fitting routine of LCModel (http://www.lcmodel.com), which computes in vivo spectra as a linear combination of complete model spectra of specific in vitro metabolites predefined in basis sets. These basis sets included spectral data of alanine, aspartate, creatine, glucose, γ-aminobutyric acid, glutamine, glutamate, scylo-inositol, myo-inositol, N-acetylaspartate, N-acetyl aspartylglutamate, glycerylphosphocholine, phosphocholine, guanidinoacetate, lipids, and macromolecules. Representative spectral data fits for normo-, de-, and rehydration are shown in Fig 1. For fitting macromolecule resonances, LCModel integrates a priori knowledge on macromolecules on the basis of metabolite-nulled spectra at 3T (ie, macromolecule models are simulated).

On the basis of the principle of reciprocity, differences in coil load were corrected for by dividing the measured metabolite resonances by the required voltage amplitude to obtain maximum signal.

**FIG 1.** Representative 1H-MR spectroscopy spectral data fits. The voxel was placed in the left paracentral lobule. Spectral data fits (upper row) and subtracted spectra (lower row) for visualization of metabolite resonances at normo- (black line, N), de- (blue line, D), and rehydration (red line, R). tCr indicates total creatine, ml, myo-inositol; tCho, total choline; NAA + NAAG, N-acetylaspartate and N-acetyl aspartylglutamate; Glx, glutamine + glutamate.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Normohyd</th>
<th>Dehyd</th>
<th>Rehyd</th>
</tr>
</thead>
<tbody>
<tr>
<td>NAAG (mM)</td>
<td>0.04 ± 0.01</td>
<td>0.03 ± 0.01</td>
<td>0.05 ± 0.02</td>
</tr>
<tr>
<td>Glu (mM)</td>
<td>2.4 ± 0.2</td>
<td>2.6 ± 0.3</td>
<td>2.4 ± 0.2</td>
</tr>
<tr>
<td>GABA (mM)</td>
<td>0.3 ± 0.01</td>
<td>0.3 ± 0.01</td>
<td>0.3 ± 0.01</td>
</tr>
<tr>
<td>Gly (mM)</td>
<td>0.3 ± 0.01</td>
<td>0.3 ± 0.01</td>
<td>0.3 ± 0.01</td>
</tr>
<tr>
<td>Cr (mM)</td>
<td>2.0 ± 0.1</td>
<td>2.0 ± 0.1</td>
<td>2.0 ± 0.1</td>
</tr>
<tr>
<td>Cho (mM)</td>
<td>3.0 ± 0.2</td>
<td>3.0 ± 0.2</td>
<td>3.0 ± 0.2</td>
</tr>
<tr>
<td>Myo (mM)</td>
<td>0.5 ± 0.01</td>
<td>0.5 ± 0.01</td>
<td>0.5 ± 0.01</td>
</tr>
<tr>
<td>NAA (mM)</td>
<td>1.5 ± 0.1</td>
<td>1.5 ± 0.1</td>
<td>1.5 ± 0.1</td>
</tr>
<tr>
<td>NAA + NAAG (mM)</td>
<td>3.0 ± 0.2</td>
<td>3.0 ± 0.2</td>
<td>3.0 ± 0.2</td>
</tr>
</tbody>
</table>

Criteria for assessment of spectral quality were the following: 1) the signal-to-noise ratio (defined by the ratio of the maximum in the spectrum-minus-baseline over the analysis window to twice the root-mean-square residuals, 2) the line width (roughly estimated by using the full width at half maximum), 3) the distribution of residuals, and 4) the SD. In the LCMedal, SD estimates are Cramer-Rao lower bounds. Cramer-Rao lower bounds of >50% indicate that the metabolite concentration may range from zero to twice the estimated concentration. In this study, we selected for further analysis only spectral data meeting the following: 1) a signal-to-noise ratio of >4; 2) full width at half maximum of ≤0.07 ppm; 3) randomly distributed residuals of ~0; and 4) Cramer-Rao lower bounds of <25%.
Quantitative T1-Mapping. Quantitative T1-relaxation time maps were estimated on a voxelwise basis according to the following formula:

\[
T1_{c,j,k} = \frac{TR}{\ln[(\sin(FA_1) \times \cos(FA_2) - Q_{j,k} \times \sin(FA_2)) \\
\times \cos(FA_1))/(\sin(FA_1) - Q_{j,k} \times \sin(FA_2))]},
\]

where \(T1_{c,j,k}\) is the T1-value, and \(Q_{j,k}\) represents the signal intensity ratio for the voxel (j,k). For this purpose, the 2 FLASH acquisitions of the 2 flip angles (\(FA_1, FA_2\)) were coregistered by using mri_robust_template (part of FreeSurfer), and the upper and lower slices were discarded. To obtain separate T1-estimates for white and cortical as well as subcortical gray matter, we rigidly coregistered the segmentations/parcellations of FreeSurfer to this individual template space (by using mri_robust_register, also part of FreeSurfer), slightly eroded and applied as masks for extracting corresponding T1-relaxation times.

Statistical Analyses
All statistical analyses were performed in SPSS Statistics (Version 22; IBM Armonk, New York). On the basis of the assumption that organic osmolutes are involved in cerebral fluid regulation, myo-inositol, creatine, glutamine, glutamine + glutamate, and N-acetyl-aspartate and N-acetyl-aspartyl-glutamate were analyzed for changes on de- and rehydration. A repeated measures ANOVA was applied to the spectroscopic estimates of these metabolites, brain tissue fluid, and serum parameters to test for experimental changes with time (ie, different levels of hydration) by using a significance level \(P \leq .05\), which corresponds to a type I error probability of \(\leq 5\%\). Thus, in the repeated measures ANOVA, the metabolite signal was the dependent measure, and time (session/level of hydration), the explanatory factor.

Post hoc \(t\) test comparisons were used to assess changes between the different pairs of longitudinal time points. On the basis of physiologically appropriate assumptions about the directionality of change (ie, that osmolality should increase while brain-water resonances and volume should decrease on dehydration and vice versa on rehydration), corresponding \(P\) values were estimated 1-sided. All statistical inferences were performed as paired tests.

The Pearson linear correlation coefficient \(r\) was used to assess 1-sided associations among brain tissue fluid, serum parameters, and morphometry data. \(P\) values were corrected for multiple comparisons (Bonferroni). Linear regression plots were generated to visualize significant associations among parameters.

Longitudinal morphometric changes of cortical surface boundaries were analyzed by using a Linear Mixed Effects Matlab toolbox (http://www.mathworks.com/help/stats/linear-mixed-effects-models.html). A piece-wise linear model was fitted to the data to detect changes on de- and rehydration.

RESULTS
Blood/Serum Parameters
On dehydration, the hematocrit and osmolality (OSMserum) measures clearly indicated loss of body fluid of all participants (Table). The concentrations of K+serum (\(P = .00\)) and serum glucose (\(P = .01\)), HCT (\(P = .00\)), and OSMserum (\(P = .03\)) changed appropriately on de- and rehydration, whereas changes of sodium and urea were insignificant.

None of the individuals examined met the criteria for acute hypernatremia (ie, mean sodium levels remained below 145 mmol/L and osmolality remained below 300 mOsm/kg for all participants. Thus, blood/serum parameters indicated subclinical dehydration consistent with 12 hours without drinking overnight. No participant reported symptoms other than fatigue and thirst or exhibited clinical signs of dehydration.

\(^1\)H-MR Spectroscopy
Brain Tissue Fluid. Normalized resonances of tissue fluid content of H2Obrain decreased on dehydration by 1.63 ± 2.84% and steadily increased by 0.43 ± 0.92% at the end rehydration (\(F = 2.49, P = .00\)) (Fig 2). Post hoc tests demonstrated a significant H2Obrain decrease from normo- to dehydration and significant increases during rehydration (On-line Table 1). Resonance changes of brain tissue fluid were associated with volume changes of the whole brain (\(r = 0.22, P = .02\)), cerebral white matter (\(r = 0.34, P = .02\)), and the hypothalamus/thalamus (\(r = 0.18, P = .05\)) (On-line Table 2).

Brain Osmolutes. Representative \(^1\)H-MR spectroscopy spectral data fits for normo-, de-, and rehydration are shown in Fig 1 (upper row); subtracted spectral fits (lower row) represent the respective signal changes. Repeated measurements ANOVA demonstrated no significant changes in myo-inositol, creatine, glutamine+glutamate, and N-acetylaspartate and N-acetyl-aspartyl-glutamate in subjects on 12 hours of dehydration and during 60 minutes of rehydration.
MR Morphometry

Volumetry. Repeated measures statistical analyses revealed longitudinal changes of the whole brain, cortical gray and white matter volume, and the subcortical hypothalamus/thalamus volume on de- and rehydration ($P = .00$) (Fig 3). Specifically, mean volumes of the whole brain, cerebral cortex, white matter, and hypothalamus/thalamus decreased on dehydration by 0.36 ± 0.68%, 0.19 ± 1.28%, 0.59 ± 0.86%, and 0.30 ± 1.80% and increased on rehydration by 0.87 ± 0.57%, 1.50 ± 1.03%, 0.23 ± 0.82%, and 1.11 ± 1.28%, respectively (Fig 3). Post hoc tests revealed significant volume changes for the whole brain, cerebral cortex, and hypothalamus/thalamus during dehydration (between normo- and rehydration) and for white matter on dehydration (On-line Table 3). Except for white matter, the magnitude of volume changes detected from normo- to dehydration was lower than that from de- to rehydration, while the associated variance was generally higher.

Correlation analyses revealed associations between the appropriate changes of the whole-brain volume and H$_2$O$_{\text{brain}}$ ($r = 0.22$, $P = .02$) (On-line Fig 1A), of the whole-brain volume and hematocrit values ($r = -0.31$, $P = .00$) (On-line Fig 1B), of the cortical gray matter volume and HCT ($r = -0.22$, $P = .02$) (On-line Fig 1C), of the cerebral white matter volume and H$_2$O$_{\text{brain}}$ ($r = 0.34$, $P = .00$) (On-line Fig 1D), of the cerebral white matter volume and HCT ($r = -0.31$, $P = .00$) (On-line Fig 1E), of the hypothalamus/thalamus volume and H$_2$O$_{\text{brain}}$ ($r = 0.18$, $P = .05$) (On-line Fig 1F), and of the hypothalamus/thalamus volume and OSM$_{\text{serum}}$ values ($r = -0.18$, $P = .04$) (On-line Fig 1G) (On-line Table 2).

Cortical Thickness and Subcortical Surface Changes. Cortical thinning prevailed on dehydration, and cortical thickening, on rehydration. These changes of cortical thickness, as averaged across subjects, were not uniformly distributed over the cerebral surface (Fig 4A). Associated error probabilities (Fig 4B) revealed a similar consistent pattern and largely excluded changes to the opposite (ie, thickening on dehydration and thinning on rehydration), which presumably reflect false-positive detections. Error probabilities ensure that the detected cortical thickness changes are not driven by outliers and accompanied by increased variance measures across subjects. Some areas attained high significance levels (ie, low error probabilities) (Fig 4B), despite relatively lower mean cortical thickness change (Fig 4A), due to low variability of the changes across subjects. The patterns of cortical thinning on dehydration and thickening on rehydration approximately mirrored each other and appeared quite similar (ie, symmetric, for the 2 hemispheres).

Quantitative T1-Relaxation Times

Mean T1-relaxation times during normo-, de-, and rehydration were 1339 ± 86.92 ms, 1314 ± 70.26 ms, and 1321 ± 75.28 ms for the cerebral cortex; 1291 ± 68.07 ms, 1285 ± 66.96 ms, and 1295 ± 68.27 ms for the hypothalamus/thalamus; and 834.52 ± 58.11 ms, 818.15 ± 42.58 ms, and 820.36 ± 46.18 ms for white matter (Fig 5). Statistical analyses revealed no longitudinal differences in T1-relaxation times of the cerebral cortex ($P = .07$), hypothalamus/thalamus ($P = .19$), and white matter ($P = .09$) on de- and rehydration.

DISCUSSION

In this study, for the first time, the effect of hyper- and hyposmolality on brain tissue fluid, brain volume, and cortical thickness was monitored in vivo. These findings represent cellular changes of the volume regulatory mechanisms at a macroscopic level.

Hypernatremia is associated with brain-water loss and corresponding volume reduction, whereas hyponatremia is related to brain-water accumulation and associated volume increase. Both conditions have been extensively studied in animal experiments and reflect osmoadaptive responses of the cell to fluid imbalances. Acute hypernatremia is defined as a plasma sodium concentration above 150 mmol/L, and acute hyponatremia, as a plasma sodium concentration below 120 mmol/L, developing during 24–48 hours. To that end, a mean plasma sodium con-
The concentration of 138.5 mmol/L on dehydration and 136.9 mmol/L after rehydration as presented in our study is considered subtle. In animal models, induced hypernatremia with 200 mmol/L led to a decrease of brain-water up to 14%, whereas a rapidly evoked (2 hours) hyponatremia with 119 mmol/L resulted in a 16% increase of brain-water. The rate at which osmolar stress develops defines whether the osmoadaptive capacities of the brain are exceeded. Therefore, it is plausible that a slowly induced (48 hours) hyponatremia with 107 mmol/L revealed no changes in brain-water.28

In the study here, 12 hours of thirsting elevated serum osmolality by 0.67% and resulted in 1.63% reduction of \( H_2O_{brain} \). One hour of slow continuous oral fluid intake lowered serum osmolality by 0.96% and led to a 0.43% increase in \( H_2O_{brain} \). This finding supports, in principle and within limits, the sensitivity and reliability of the \( H_2O_{brain} \) signal, which is, especially with receive-only multichannel phased array coils, commonly used for water scaling in \(^1\)H-MR spectroscopy.19,20,29

On de- and rehydration, changes of the global brain volume, gray and white matter volume, hypothalamus/thalamus volume, and cortical thickness were observed, which were consistent with the processes of cell shrinkage and swelling, respectively, and are in line with findings in previous studies.10-13 Global brain volume changes on dehydration (0.36%) and rehydration (0.87%) are in excellent agreement with those in a previous study,13 which reported volume changes of the same magnitude after 16 hours of thirsting (0.55%) and subsequent rehydration (0.72%). Previous studies of Dickson et al,11 Kempton et al,10 and Watson et al,12 which induced dehydration by thermal exercises, failed to demonstrate brain volume changes.

Freund et al15 demonstrated a reversible volume reduction of cerebral gray matter (6%) after 2 months of daily running during an ultramarathon, which is in line with the reversible cortical thinning (0.19%) on dehydration reported in our study. On the other hand, a recent study with low sample size14 failed to reveal thinning of the cerebral cortex and shrinking of the hypothalamus/thalamus, which are reversed on rehydration. Reversible cortical thinning by short-term dehydration is not uniformly distributed over the cerebral surface (Fig 4) (ie, the brain does not seem to simply shrink by global scaling during dehydration). Instead, detected changes of cortical thickness reveal some but no exclusive overlap with the following: 1) known areas of pronounced cortical thickness30; and 2) atrophy patterns found in Alzheimer disease,31 cerebrovascular dementia,32 and eating disorders,33 for example. Also, more generally, brain volume changes due to osmotic stress are in/near the magnitude of

**FIG 4.** Cortical thickness analysis. Local changes of cortical thickness (A, red-to-yellow: for thinning on dehydration; blue-to-light blue: for thickening on rehydration). Dehydration primarily induces cortical thinning (upper row), which reverses on rehydration (bottom row). Note that these prevailing changes are not uniformly distributed over the cerebral surface. Changes on the mesial surface (not shown) were slightly less pronounced but similar. The corresponding statistical significance (B, red-to-yellow: for thinning on dehydration; blue-to-light blue: for thickening on rehydration) is expressed by increasingly lower false-positive probabilities across subjects.

**FIG 5.** T1-relaxometry. Quantitative T1-relaxation times of the cerebral cortex (upper row), the subcortical gray (middle row), and white matter (bottom row) during normo-, de-, and rehydration. Statistical analyses revealed no longitudinal changes in T1 relaxation times. Box indicates upper and lower quartiles; thick black line, median; whiskers, most extreme values of the interquartile range; crosses, outliers.
changes attributed to annual volume decrease by aging (0.2%), 34,35 Alzheimer disease (2%), Lewy body dementia (1.4%), or vascular dementia (1.9%). 36 These results emphasize that longitudinal changes and cross-sectional differences of cortical thickness in conditions accompanied by altered fluid intake (such as dementias or eating disorders) can be confounded by dehydration, which may then be mistaken for brain atrophy or even regeneration.

Whole-brain, white matter, and hypothalamus/thalamus volumes demonstrated changes related to those in brain tissue fluid. This finding is further supported by the inverse correlation between HCT and volume changes of the whole brain, cerebral cortex, and cerebral white matter. The only brain structure that demonstrated an association between its volume and serum osmolality changes was the hypothalamus/thalamus. This finding is consistent with the fact that the hypothalamus includes magnocellular neurosecretory cells of the paraventricular and supraoptic nucleus, which are able to intrinsically detect and respond to changes in osmolality. 37 The magnocellular neurosecretory cell volume has been shown to be inversely proportional to the osmolality of the extracellular fluid. 38 Moreover, in a magnocellular neurosecretory cell subset of the paraventricular and supraoptic nucleus, vasopressin is synthesized. Axons project into the neurohypophysis, where vasopressin release occurs. Thus, the association between volume changes of the hypothalamus/thalamus as found in our study may reflect the relationship between magnocellular neurosecretory cell volume and osmolality of the extracellular fluid. However, we acknowledge that magnocellular neurosecretory cells are distributed over different subcortical gray matter nuclei and cannot be directly assessed by MR imaging at 3T.

No statistically significant changes of cerebral osmolytes on dehydration or after rehydration were observed. This finding may be due to these changes falling below the between- and within-session reproducibility limits of the spectroscopy measurements. Note that the between-session variance exceeded the within-session variance across subjects for all morphometric and spectroscopic measures studied (Figs 2, 3, and 5), which presumably results from repositioning and reshimming. Alternatively, 12 hours of mild dehydration may not produce changes in organic osmolytes large enough to be detectable by MR spectroscopy—at least not in all individuals—that is, osmolytes may already act as osmotic active substrates in some individuals but not in others. Although the resulting metabolite data might demonstrate plausible kinetics of organic osmolytes in a subset of individuals, there is a high variance in the data across subjects, which emphasizes such interindividual differences. In this study, mild dehydration was associated with a 0.67% increase in serum osmolality, and rehydration resulted in a 0.29% decrease in serum osmolality. For comparison, Videen et al 39 reported cerebral osmolyte changes (reduction of myo-inositol, choline, total creatine, and N-acetylaspartate) on hyponatremia in patients with pituitary tumors, congestive heart failure, or syndrome of inappropriate antidiuretic hormone secretion associated with an average 12.98% reduction in serum osmolality. Lien et al 40 examined severe hyponatremia in rats, which led to a mean 33.55% increase in serum osmolality and also demonstrated changes in organic osmolytes (increase of myo-inositol, choline, phosphocreatine, glutamine, and glutamate) in brain tissue.

Limitations of this study are methodologically inherent because assigning resonances of 1H-MR spectroscopy data to metabolites, for example, is itself error-prone. Data quality and selection of an adequate fitting routine are therefore essential.

At least in theory, morphometric changes may also result from purely physical changes in tissue relaxation times and contrast induced by fluid changes influencing tissue segmentation. However, there were no changes in T1-relaxation times of the cerebral cortex and the subcortical gray and the white matter among normo-, de-, and rehydration, while the average T1-values themselves were in excellent agreement with previous data. 41 Also, the correlations between morphometry measures and serum parameters support physiologic effects. Moreover, whether physiologic and/or physical, the key message is that hydration levels constitute an important confound for morphometric studies, and this is the first study demonstrating relevant changes of cortical thickness related to subclinical dehydration and rehydration. Brain tissue fluid resonances, rapidly assessed by unsuppressed 1H-MR spectroscopy, are suitable for water scaling and, along with serum parameters, also correlate with whole-brain, white matter, and hypothalamus/thalamus volume changes.

CONCLUSIONS

This study suggests the following: 1) Morphometric brain changes during de- and rehydration match the concept of osmotically induced cell volume changes, and 2) the cerebral cortex, white matter, and the hypothalamus/thalamus are affected by these changes. Our results emphasize that it is essential to control for hydration levels in any study on cerebral morphometry or metabolism to avoid confounding the findings. This is especially important because hydration levels can systematically differ across population samples and longitudinal follow-up, in particular for patients with dementias, eating disorders, and other conditions accompanied by even slightly altered fluid intake.

ACKNOWLEDGMENTS

We thank Professor Andreas Draguhn (Institute of Physiology and Pathophysiology, Division of Neuro- and Sensory Physiology, University of Heidelberg, Germany) for critical reading of the manuscript and very helpful comments.

Disclosures: Martin Reuter—RELATED: Grant: National Institutes of Health, Grant K25 CA188632.* Martin Bendszus—UNRELATED: Consultancy: Codman; Payment for Lectures (including service on Speakers Bureaus): Novartis, Guerbet, Codman, Roche. *Money paid to the institution.

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