Supporting Information

Lipid Removal in Deuterium Metabolic Imaging (DMI) using Spatial Prior Knowledge.

Robin A. de Graaf 1,2, Yanning Liu 2, Zachary A. Corbin 3, Henk M. De Feyter 1

Departments of Radiology and Biomedical Imaging 1, Biomedical Engineering 2 and Neurology 3
Magnetic Resonance Research Center
Yale University School of Medicine
New Haven, Connecticut, USA

Address correspondence to:
Robin A. de Graaf, Ph.D.
Magnetic Resonance Research Center
Department of Radiology and Biomedical Imaging
Yale University School of Medicine
300 Cedar Street, P.O. Box 208043
New Haven, CT 06520-8043, USA
Tel: (203) 785-6203
Fax: (203) 785-6643
E-mail: robin.degraaf@yale.edu
Figure S1 – Examples of brain and skull constellations used in simulations. (A) Brain and skull ROIs used in simulations on lipid suppression and metabolite retention. Brain ROIs of healthy volunteers were segmented into gray matter (GM), white matter (WM) and cerebrospinal fluid (CSF) after which a pathology ROI was placed at a random location. Each simulation was accompanied by randomly generated (B) $B_0$ and (C) relative $B_1$ maps based on ranges established in vivo (e.g., Fig. 1E/F). Anatomy-based ROIs were defined on five subjects, with ten permutations in pathology location and size and $B_0/B_1$ distribution per subject, for a total of 50 datasets.
Figure S2 – Lipid suppression and metabolite retention for different skull ROI divisions. (A) Lipid suppression and (B) metabolite retention under different scenarios including \( N_{\text{skull}} = 1 \) without (column I) and with (column II) \( B_0 \) and \( B_1 \) compensation, \( N_{\text{skull}} = 20-28 \) (column III), 29-36 (column IV), 37-48 (columns V and VI) without (columns III-V) and with \( B_0 \) and \( B_1 \) compensation (column VI). Panels (A) and (B) are identical to Fig. 2G/H. (C) Metabolite retention for the separate brain ROIs (GM, WM, CSF, and pathology) under identical scenarios as in (A/B). Metabolite retention in separate brain ROIs (C) closely follow that across the entire brain (B).
Figure S3 – Absolute-valued SRF contributions of varying number of skull ROIs to a single brain ROI. All calculations used the brain/skull constellation shown in Fig. 2 with the four brain compartments (GM, WM, CSF, pathology) combined into a single brain ROI. (A-D) Example skull subdivisions for \( N_{\text{skull}} = 1 \) (A), 10 (B), 30 (C) and 50 (D). The first (dark red) column represents the SRF contribution of the brain SRF to the brain ROI which is, by definition, 1.0 and is shown for intensity reference. For a coarse subdivision of the skull ROI (\( N_{\text{skull}} < 30-35 \)), the SRF contribution of each skull ROI is low, in agreement with Fig. 2K. Note that summation over the real-valued SRF will produce a zero SRF contribution for all skull ROIs for any \( N_{\text{skull}} \). The small absolute-valued SRF contribution seen for \( N_{\text{skull}} < 30-35 \) means that phase cancellation of signal across the brain compartment is only a minor contributor to the overall localization performance. While the skull SRFs are well-behaved, the small number of skull ROIs only provide modest immunity to \( B_0/B_1 \) heterogeneity (Fig. 2G, column III). Finer subdivision of the skull ROI (\( N_{\text{skull}} > 35 \)) provides improved lipid suppression (Fig. 2G, columns IV and V), at the cost of greatly increased skull SRF contributions to the brain ROI (Fig. 2N). Even though phase cancellation provides perfect localization under homogeneous conditions, intra-compartment heterogeneity will lead to incomplete phase cancelation and to increased signal variation in the brain compartment (Fig. 2H, column V). The sharp transition from low to high SRF contributions at \( N_{\text{skull}} \sim 35 \) is similar for all 50 simulated brain/skull configurations.