Optimized Processing Of Quantitative Susceptibility Mapping-Based Gadolinium Perfusion MRI: Correction Of Bulk Susceptibility Effects And Comparison Of Arterial Input Function Selection From ΔR₂* and QSM Data

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Target Audience. Researchers and clinicians working with MR perfusion imaging. Purpose. Analysis of dynamic MRI during bolus injection of Gd contrast agent is traditionally performed by repeated measurement of brain signal relaxation times (usually T2*) to estimate Gd concentration, however a nonlinear relationship between ΔR₂* and Gd tissue concentration has been reported12. Quantitative Susceptibility Mapping (QSM) estimates the tissue susceptibility distribution Δγ(r) from the MR phase data3,4. Δγ is expected to represent a different contrast mechanism and to be less affected by nonlinearities in the estimation of the Gd concentration (molar susceptibility χm). In our previous work5, we have demonstrated the feasibility of dynamic QSM data Δγ(r,t) reconstruction for the measurement of cerebral blood flow (CBF). This abstract presents a refined QSM-based analysis scheme which addresses bulk Gd effects, and compares QSM and ΔR₂*-based analyses. The influence of arterial input function (AIF) selection from both the QSM data and the ΔR₂* data is also investigated. Methods. Experiment. A 68 year-old male patient without perfusion deficit was scanned at 3.0T (Philips Healthcare) as part of his stroke workup for symptoms of blurred vision. Informed consent and IRB approval were obtained; Scan parameters. 20 slices, 80 dynamics, multi-slice 2D gradient echo EPI, FOV=240x240 mm², 256x256 matrix, 5mm slice thickness, TR/TE 1500/40 msec, 0.2 mmol/kg Gd bolus. Analysis. Perfusion analysis utilized the magnitude data to determine ΔR₂*(t)6. QSM processing included Laplacian phase unwrapping7, frequency map calculation f(t), background field removal (dipole fitting) and susceptibility calculation (LSOR method)7,8. Compared to the previously presented1 data (to effectively avoid artificial susceptibility decrease in brain voxels during bolus passage), the Δγ(t) maps were corrected by adding the average background susceptibility change estimated from the dipole fitting analysis before perfusion processing. This step is necessary to account for bulk Gd effects which would otherwise be removed with the background. Perfusion parameters (CBF, CBV and MTT) were calculated using standard perfusion processing6 with Tikhonov regularization9, using both ΔR₂*(t) and Δγ(t) maps. Two semi-automatically detected arterial input functions (AIF)10 were determined: a QSM-derived AIF used for QSM processing only, and a DSC-derived AIF applied to both analyses. Due to dynamic range and aliasing, the QSM-derived AIF shape was determined from voxels adjacent to the MCA and scaled to the expected peak concentration by the use of a scaling factor determined from the equilibrium concentration. Quantitative analysis used a transverse relaxivity of Gd r₂=5.2 L•mol⁻¹•sec⁻¹ and molar susceptibility χm=3.290×10⁻³ L•mol⁻¹ for quantitative conversion2,11. Results. Fig. 1 depicts the perfusion parameter maps for CBF, CBV and MTT determined with ΔR₂* and QSM (for both AIF choices). QSM-derived perfusion shows comparable data quality and demonstrates good general qualitative and quantitative agreement to DSC data. While CBF results are most similar, both CBV and MTT show decreased values in WM for QSM-derived maps. Fig. 2 depicts the DSC-derived AIF (Fig.2a) and the QSM-derived AIF (Fig.2b). Both were converted into units of mM of Gd. QSM and ΔR₂*-derived AIF are of comparable quality and show good quantitative agreement. Discussion. An improved processing scheme of QSM perfusion data achieves comparable data quality to DSC. QSM may have decreased sensitivity to slowly perfused WM due to SNR limitations (currently, respiratory and extracranial Gd related phase changes cannot be fully eliminated). Local differences in the observed WM and GM perfusion may reflect different microvascular sensitivities of both methods, as the underlying contrast mechanism of ΔR₂* exploits the dephasing related signal amplitude loss, while QSM is based on the average detected local resonance frequency. Thus, QSM may add complementary information to MR perfusion data, while both ΔR₂* and QSM perfusion are characterized by a sensitivity dominated by large vessels. Despite SNR limitations, the QSM-derived AIF shape and peak is very similar to the ΔR₂*-derived AIF. QSM-AIF calculation currently requires a scaling factor estimated from the steady state susceptibility change. Conclusion. A refined processing scheme for QSM perfusion analysis identifies and addresses bulk Gd susceptibility changes and leads to good agreement to DSC perfusion parameters. Separation of background from contrast-induced phase changes remains a challenge. Some of the QSM perfusion information may be complementary to ΔR₂*. Accurate AIF curves can be determined from the QSM data, and QSM- and ΔR₂*-derived AIF yield comparable perfusion results. References. 1. F Calamante, et al. Magn Reson Med. 58:544-53, 2007. 2. MJ van Osch, et al. Magn Reson Med. 49:1067-76, 2003. 3. W Li, et al. Neuroimage. 55:1645-56, 2011. 4. F Schweser, et al. Neuroimage. 54:2789-807, 2011. 5. D Bonekamp, et al. Proc Int Soc Mag Reson Med, pg. 1937, 2011. 6. L Ostergaard, et al. Magn Reson Med. 36:715-25, 1996. 7. L de Rochefort, et al. Magn Reson Med. 63:194-206, 2010. 8. S Wharton, et al. Magn Reson Med. 63:1292-304, 2010. 9. S Sourbourn, et al. Phys Med Biol. 49:3307-24, 2004. 10. L Knutsson, et al. 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Figure 1

Figure 2