Introduction: Gadobenate dimeglumine is a Gd-based chelate that is widely used for contrast enhanced magnetic resonance angiography (CE-MRA) procedures. Although not a dedicated intravascular agent, it interacts weakly and transiently with serum albumin resulting in increased longitudinal relaxivity ($r_1$) relative to conventional Gd-based contrast agents. Recently, Anzidei et al. (1) have shown that the increased $r_1$ renders gadobenate dimeglumine suitable for Steady State (SS) CE-MRA and that the additional diagnostic information available on combined first passage (FP) and SS CE-MRA is potentially greater than that on FP alone for both carotid and peripheral CE-MRA. As yet, however, few studies have been performed to optimize the acquisition protocols for CE-MRA with gadobenate dimeglumine. The present investigation aimed to define the optimal flip angle (FA) for gadobenate dimeglumine-enhanced SS MRA and was performed in three distinct phases: 1) in vitro optimization of an inversion recovery protocols for dynamical monitoring of the longitudinal relaxation rate $r_1$; 2) in vivo dynamical evaluation of $r_1$ in blood after administration of gadobenate dimeglumine and determination, for different repetition times (TR), of the optimal flip angle for SS MRA; 3) assessment of the benefits of using a calibrated Ernst Angle in SS MRA of carotid arteries.

Methods: In vitro optimization: Phantom studies were performed at 1.5 T using a clinical whole body MR scanner equipped with a transmitting body coil and a receiving head coil. A multi-tubes test phantom was prepared by mixing in different concentration a commercial formulation of 0.5 mol/L gadobenate dimeglumine with a physiological saline solution. The reference standard for the $r_1$ evaluation was a spin echo (SE) sequence, in which TR was varied between 20 and 10,000 ms to monitor the exponential recovery of the longitudinal magnetization. The inversion recovery fast low angle shot (IR-FLASH) sequence to be subsequently exploited in vivo has been tested against the SE for different combinations of flip angle (FA, between 6° and 14°) and inversion time (TI, between 134 and 9,000 ms). In vivo studies: 25 patients (17 male, 8 female; age: 54 ± 3 years; range: 48 – 70 years) scheduled for a contrast-enhanced MR examination for non-vascular indications were enrolled after approval of the Local Institutional Committee of Medical Ethics. All subjects provided written informed consent after having been informed of the potential benefits and contraindications of CE-MRA. On five patients, the optimized IR-FLASH sequence was used to monitor the time evolution of blood $r_1$ at 1.5 T after administration of gadobenate dimeglumine at 0.1 mmol/kg bodyweight. An optimal FA for angiographic measurement was thereafter derived from the Ernst equation based on experimental values of $r_1$ previously determined in vivo. Steady state CE-MRA examinations of carotid arteries were performed in the remaining 20 subjects by means of a 3D gradient echo sequence in order to evaluate the improved contrast achieved after optimization of the FA for maximal blood signal enhancement.

Results: An ultra-fast IR-FLASH sequence with FA=8° and a properly defined set of TI values was shown to give in vitro $r_1$ determinations in good agreement with those obtained by a routine variable-TR SE sequence. The use of this ultra-fast IR-FLASH sequence in vivo allowed the blood signal behavior in the carotid arteries after gadobenate dimeglumine administration to be monitored. $r_1$ was found to decrease exponentially from 8.7±0.96 s$^{-1}$ at 30 sec post injection to 3.8±0.24 s$^{-1}$ at 10 min post injection, in agreement with theoretical predictions based on gadobenate dimeglumine pharmacokinetic data (2) and in vitro $r_1$ relaxivity (3,4). Optimal flip angles, at different TR, for SS angiography with gadobenate dimeglumine, derived according to Ernst equation from our $r_1$ experimental data are displayed in Fig. 1. Significantly higher blood Signal to Noise Ratio (SNR) was achieved on SS images of carotid arteries acquired using a 3D spoiled gradient echo sequence with TR = 7.5 ms and FA=18° than on corresponding images acquired with a FA=35° as used typically for intravascular blood pool contrast agents (52.5 ± 8.3 vs. 29.5 ± 6.0; p<0.05, Mann Whitney U test), as displayed in Fig. 2.