Target audience: Radiologists and physicists with an interest in functional MRI of the kidney and small animal imaging.

Purpose: To investigate whether multiparametric functional MRI allows detection of acute renal allograft rejection and to compare MRI parameters with renal histopathology and the composition of cell infiltrates of renal tissue.

Methods: An acute renal allograft rejection was induced by allogenic kidney transplantation (ktx) of C57Bl/6-kidneys to Balb/c-mice (n=6). Animals after isogenic ktx (C57Bl/6-kidneys to C57Bl/6-mice) were used as controls (n=5; Figure 1). MRI was performed three weeks after ktx using a 7 Tesla magnet (Bruker, Pharmascan). Renal perfusion was quantified using a flow sensitive alternating inversion recovery (FAIR) EPI ASL sequence (13 TI = 30-8000 ms)². In order to evaluate tissue edema, maps of T1- and T2-relaxation time were calculated. In addition, apparent diffusion coefficients (ADC) were determined from an echoplanar diffusion-weighted sequence (7 b-values = 0-700 s/mm²) using a monoexponential fit³. After the MRI, animals were sacrificed and histological changes of renal tissue were evaluated according to Banff criteria. Furthermore, the composition of infiltrating cells of renal tissue was assessed by FACS-analysis. Differences between groups of allogenic and isogenic ktx were evaluated using unpaired t-tests and the correlation of functional MRI parameters with the percentage of T-cell infiltrates was determined (Pearson). Values are given as mean±SEM.

Results: Animals after allogenic ktx developed an acute T-cell-mediated rejection according to Banff criteria, whereas renal histology after isogenic ktx was unremarkable. The percentage of infiltrating T-cells was significantly higher after allogenic (38.8±4.0%) than after isogenic ktx (5.5±2.2%; p<0.001). Renal perfusion was significantly impaired in animals with an acute rejection (allogenic ktx) compared to the control group (56±7 vs. 293±44 ml/(min*100g); p<0.001; Figure 2). T1- and T2-relaxation times of renal tissue were increased after allogenic ktx with most pronounced changes in the outer medulla. Here, T1 values were 1938±53 ms after allogenic and only 1349±27 ms after isogenic ktx (p<0.001). T2 values were 60.1±2.0 ms and 45.7±1.2 ms, respectively (p<0.001). ADC values were significantly reduced after allogenic (1.39±0.14*10⁻³ mm²/s) compared to isogenic ktx (1.83±0.05*10⁻³ mm²/s; p<0.05). The percentage of T-cell infiltrates negatively correlated with renal perfusion (r=-0.84) and ADC (r=-0.78) and positively with T1- (r=0.97) and T2-relaxation times of renal tissue (r=0.92).

Discussion: Functional MRI allows detection of acute, T-cell-mediated renal allograft rejection. This is associated with impairment of renal perfusion, increase of T1- and T2-values, interpreted as tissue edema due to inflammation, and ADC reduction due to cellular infiltration. MRI parameters correlate with the percentage of T-cell infiltrates. Thus, multiparametric functional MRI may improve non-invasive diagnosis of renal allograft rejection.

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