In vivo brain connectivity: optimization of manganese enhanced MRI for neuronal tract tracing.

Santiago Canals, Michael Beyerlein, Anna L Keller, Yusuke Murayama and Nikos K Logothetis

Physiology of Cognitive processes, Max Planck Institute for Biological Cybernetics, Spemannstrasse 38, Tübingen, Germany Email: santiago.canals@tuebingen.mpg.de

One of the main problems in systems biology is to obtain information on signal processing between interconnected groups of neurons in highly distributed networks. The recently introduced technique of manganese (Mn^{2+}) enhanced MRI (MEMRI) to study neuronal connectivity *in vivo* opens the possibility to these studies. However, several drawbacks exist that challenge its applicability. High Mn^{2+} concentrations produce cytotoxic effects that can perturb the circuits under study. In the other hand, the MR signal is proportional to the Mn^{2+} concentration in tissue and thus, significant amounts of Mn^{2+} are required to produce detectable contrast and reliable connectivity maps.

Here we attempt to optimize the MEMRI technique by preventing toxicity and improving the quality and extension of the obtained connectivity maps.

The somatosensory cortex of male SD rats was stereotaxically injected with different Mn^{2+} -containing solutions. Total amount of injected Mn^{2+} ranged between 1 and 16 nmol and the injected volumes between 10 and 80 nL. Osmolarity and pH effects were investigated injecting pH buffered solutions of Mn^{2+} (pH 7.3 in Tris-HCl buffer vs. 5.5 in H₂O) at different concentration (0.05, 0.1 and 0.8 M MnCl₂). Same amounts of Mn^{2+} (8nmol) delivered to the tissue at different infusion rates were also compared. Following the injection, T₁-weighted MR imaging (250 mm isotropic resolution) was performed in a 7T scanner at different time points. Fifteen days after the injection animals were sacrificed and brains processed for histology. Nissl staining as well as GFAP and NeuN immunohistochemistry (selective staining for astrocytes and neurons, respectively) were performed in the brain sections to examine cellular toxicity.

All injections produced connectivity maps consistent with the known anterograde projections of SI cortex based on classical neuronal tract-tracing techniques. Our results show that pH buffered solution improve the effectiveness of MEMRI, increasing T₁ contrast in the projection sites. In addition, injections of pH buffered and isotonic solutions of 50 and 100 mM MnCl₂ yielded more extensive connectivity maps, in particular, ipsiand contra-lateral corticocortical connections were evident in all animal injected with those solutions but not with the more usual MEMRI protocol (0.8M MnCl₂ in H₂O). Hypertonic and non-buffered solutions containing 8nmol Mn²⁺ resulted in neuronal death and astrogliosis in extensive areas around the injection point. In sharp contrast, no neuronal toxicity was observed with injections containing up to 8nmol of Mn²⁺ in isotonic solutions of up to 100 mM MnCl₂ and pH 7.3. Slow infusion rates demonstrated also to be advantageous and permitted application of larger amounts of Mn²⁺ without toxic effects, resulting in better T₁ contrast in the low density projection fields.

We conclude that refined protocols for MEMRI improve the quality and extension of connectivity maps and preserves tissue viability, assuring the application of this technique in longitudinal experiments.