Transcriptomic comparative analysis of hippocampal tissue and primary cultures after hyaluronidase treatment

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Motivation and Aim: It is well-known that traumatic injury or enzymatic digestion of brain extracellular matrix can cause seizure-like activity in primary hippocampal culture [1]. However, underlying molecular mechanism and similarity between *in vitro* and *in vivo* models of epilepsy is poorly investigated. The aim of this study was to investigate changes of gene expression and perform comparative transcriptomic analysis while modeling of epileptogenesis.

Methods and Algorithms: C57BL/6J mice were used to prepare hippocampal cell culture and intra-hippocampal hyaluronidase injection. Hyaluronic acid, which is the basis of extracellular matrix, has been removed by 75U/ml hyaluronidase. Transcriptomic analysis was performed by mouse full-genome 2-colour Agilent Microarrays. Normalization, processing and detection of differentially-expressed genes were performed by online free software – Babelomics 5.0 [2]. Functional annotation clustering was made by David bioinformatics resource [3].

Results: Hyaluronan digestion resulted in significant changes of expression of 3 gene clusters in neuronal hippocampal cultures – synaptic gene cluster, ribosomal gene cluster and mitochondrial gene cluster, while in hippocampus hyaluronidase injection resulted in changes of such functional gene clusters: synaptic gene cluster, neurogenesis gene cluster, actin-binding gene cluster, circadian rhythm cluster and ubiquitinylation gene cluster. When comparing control groups of *in vivo* and *in vitro* samples there were found 3 significant gene clusters: calcium ion-binding cluster, cluster of genes expressing EGF-domains and cortical cytoskeleton gene cluster.

Conclusion: The results obtained in this study suggest that hyaluronic acid digestion mainly affect synaptic gene cluster that indicate to similarity of *in vivo* and *in vitro* models. A wide variety of gene clusters from isolated hippocampus point to high complexity of biological processes in mice brain. Transcriptomic comparative analysis revealed strong difference between cell culture and hippocampus that should be considered when interpreting *in vitro* data.

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References

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