## Insights in Clinical and Cellular Immunology

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Brain response in some systemic immune condition-Toxicological aspects

From biomedical literature "autism disorder are involved in young patient, that we have abnormalities (Imaging, histology) in some brain areas, and a comples symptomatology. Genetic and environment can produce some unbalances in brain grow and immunitary situation is involved. Apoptotic signal contribute in brain growth and immunologic shock can unbalance the environment producing abnormalities."

We can see that some pharmacological molecules are been introduced in therapy in some brain pathologies with a specific mechanism: modulating the immune systems. We can see that some systemic immune modifications can unbalance this systems producing pharmacological effect in local place (as Brain). We can observe this phenomena like a kind of toxicity that can be deeply investigate to discover new Pharmacological strategies.

Aim of this work is to observe this kind of pathologies under a specific immune-toxicological aspect. We think that in this field are needed deeply new approach in order to adequately focus this kind of disorder. A different way to set this kind of pathologies can help in searching new pharmacological strategies.

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Dendritic cells and TNF-Related apoptosis inducing ligand (TRAIL) represent new possibilities for sepsis treatment

Sepsis refers to a generalized inflammatory response of the organism to an infection or to bacterial products in circulation, rather than the development of an infection per se. Despite recent advances in clinical practice and overall medical care, sepsis remains a great health care problem and is still the most common cause of death in critically ill patients with infection. We suppose that during the course of sepsis the expression of TRAIL in different organs correlates with acute mortality and further development of multiple organ dysfunction syndrome (MODS). It is expected that dendritic cells (DCs) might become targets for apoptotic processes in a result of elevated TRAIL expression. This hypothesis is a bias for detailed investigations for in vivo studies in animal models and for in vitro studies of septic patients.