

Fluorescence studies reveal a high-affinity interaction between the neurotoxic amyloid β -peptide and calmodulin

P01-17

I. Corbacho^I, M. Berrocal^I, K. Török^{II}, C. Gutierrez-Merino^I, **A.M. Mata^I**

^IDept. Biochemistry and Molecular Biology, Faculty of Sciences, University of Extremadura, Avda. de Elvas, s/n, 06006, Badajoz, Spain, ^{II}Molecular and Clinical Sciences Research Institute, St. George's, University of London, Cranmer Terrace, SW17 0RE, London, United Kingdom

Many works have been focused on Amyloid β -peptides ($A\beta$), a major hallmark of Alzheimer's disease (AD), and their interactions with different proteins. Although $A\beta$ neurotoxicity develops with cytosolic calcium dysregulation, our knowledge about $A\beta$ and calcium regulation and signaling is still limited. One of the proteins that plays a major multifunctional role in calcium signaling is calmodulin (CaM). In this work, using fluorescent derivatives of CaM (Badan-CaM), we show that $A\beta$ binds with high affinity to CaM. Furthermore, we have been able to fine tune that the 25-35 domain of $A\beta$ form a novel binding motif for CaM which is responsible for this high affinity interaction. The low values obtained for the dissociation constants of $A\beta$ from CaM point out that CaM is one of the cellular targets with highest affinity for neurotoxic $A\beta$ peptides. Therefore, this novel high affinity $A\beta$ -CaM interaction opens a new gateway to further understand the mechanism involved in the neurotoxic effect of $A\beta$ and also to consider the potential of calmodulin and calmodulin-derived peptides as therapeutic agents in AD. *This work has been supported by Grant BFU2014-53641-P of the Spanish Plan Nacional de I+D+I and by Grant GR15139 of the Junta de Extremadura to the Research Group BBB008, both with co-financing by the European Funds for Structural Development (FEDER).*