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Synchrotron light in the structural studies of human transferrin for its application as a drug transporter in the central nervous system

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Abstract

Human serum transferrin (Tf) is a ~80 kDa glycoprotein that transits the blood plasma, whose function is to transport iron from its absorption site towards every cell through the blood vessels. Its structure consists of two homologous lobes, an N-terminal lobe and a C-terminal lobe, which can each bind a metallic ion, mainly Fe³⁺, through a coordination bond with octahedral geometry^{1–3}. This important protein has been proposed as a potential drug transporter due to its transport mechanism across the blood-brain barrier^{4,5}. To determine the feasibility of such system, a thorough structural and bioanalytical characterization of transferrin is essential, for which synchrotron light is a powerful resource. X-ray crystallography is one of the most widely used approaches for structure determination, and several Tf models have been obtained so far^{2,3,6–8}; however, further research is necessary to generate high resolution models. Besides, a plethora of non-conventional crystallization techniques are available for crystal growth optimization, which in turn provides higher quality crystallographic data^{9–11}. Therefore, the current project is exploring several of these methods for growing high quality transferrin crystals, being counter diffusion one of the main bets. Another powerful technique is Small Angle X-ray Scattering (SAXS), which has been able to provide information for Tf regarding its stability and behavior in solution¹². Since transferrin's iron release mechanism has been shown to be pH dependent¹, SAXS analyses at different pH conditions are being performed to study its conformation states throughout the stages of its iron binding and release cycle.

About

Primary author(s) : Dr. CAMPOS-ESCAMILLA, Camila (Institute of Chemistry, National Autonomous University of Mexico); Dr. SILIQI, Dritan (Institute of Crystallography (IC), National Research Council (CNR)); Dr. GAVIRA, José (Laboratorio de Estudios Cristalográficos, Instituto Andaluz de Ciencias de la Tierra (Consejo Superior de Investigaciones Científicas-Universidad de Granada)); Dr. GONZÁLEZ-RAMÍREZ, Luis (Laboratorio de Estudios Cristalográficos, Instituto Andaluz de Ciencias de la Tierra (Consejo Superior de Investigaciones Científicas-Universidad de Granada)); Dr. MORENO, Abel (Institute of Chemistry, National Autonomous University of Mexico)

Presenter(s) : Dr. CAMPOS-ESCAMILLA, Camila (Institute of Chemistry, National Autonomous University of Mexico)

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