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Modulating the cardiotoxic behaviour of immunoglobulin light chain dimers through point mutations

Martina Maritan^a, Arianna Ambrosetti^a, Luca Oberti^a, Alberto Barbiroli^b, Luisa Diomede^c, Margherita Romeo^c, Francesca Lavatelli^d, Pietro Sormanni^e , Giovanni Palladini^d, Martino Bolognesi^a, Giampaolo Merlini^d and Stefano Ricagno^a

^aDepartment of Bioscience, University of Milan, Milan, Italy; ^bDepartment of Nutritional Science, University of Milan, Milan, Italy; ^cIRCCS-Istituto di Ricerche Farmacologiche "Mario Negri", Milano, Italy; ^dAmyloidosis Research and Treatment Center, Fondazione IRCCS Policlinico San Matteo, Pavia, Italy; ^eDepartment of Chemistry, University of Cambridge, Cambridge, UK

Background

Light chain (LC) amyloidosis (AL) is caused by the overproduction, misfolding and aggregation of immunoglobulin LCs that tend to deposit as amyloid fibrils in the cardiac tissue [1]. Nowadays, the molecular details underling LCs soluble cardiotoxicity and fibril formation remain to be fully elucidated. It has been suggested a relationship between conformational flexibility and amyloidogenicity, indicating protein thermal stability and dynamics as factors able to influence the complex processes of misfolding and aggregation [2,3]. To date, no compelling correlations have been identified between LCs cardiotoxicity and primary sequence. Here, we present a pivotal mutagenesis study in which we produce and characterize a mutated variant of the toxic H6 LC (mH6). Our purpose is to abrogate the toxicity typical of H6 yielding to a non-cardiotoxic LC and to understand the biophysical and biochemical changes underlying such loss of toxicity.

Material and methods

H6 mutant design was carried out combining the results of bioinformatics tools for protein stability predictions and sequence multi-alignment. The H6 sequence was mutated into mH6 using side directed mutagenesis. *Escherichia coli* is used to express LCs, which are then refolded and purified by chromatographic methods as previously reported [3]. For protein characterization both spectroscopic (circular dichroism and fluorescence) and structural (X-ray crystallography) techniques were applied. *In-vivo* toxicity assessment was carried out measuring *Caenorhabditis elegans*' pharynx pumping rate [4].

Results

The design of mutated H6 was performed applying an innovative method that correlates structural data with toxicity (Figure 1). A set of thermal stabilizing mutation

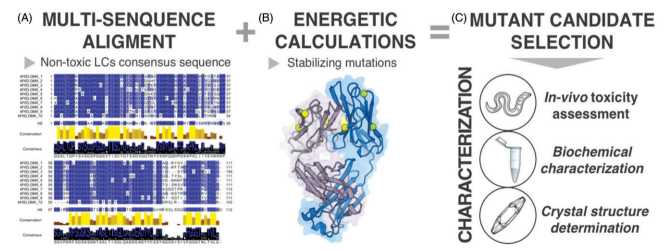


Figure 1. (A) Representative sequence multi-alignment of myeloma LCs against H6. Sequence conservation and consensus sequence are depicted by histograms. (B) Bioinformatics analysis using H6 structure as input run to identify possible mutation spots. (C) The combination of outputs from (A) and (B) lead the mutation spot selection and subsequent protein production and characterization.

candidates were found through a hybrid approach involving i) multi-sequence analysis and ii) energetic calculations. The H6 amino acid sequence was aligned against a multiple myeloma sequence database, in order to identify the conserved amino acids spots associated with nontoxic LC (Figure 1(A)). The crystal structure of H6 was analysed by bioinformatics tools that calculated the most favourable mutations to achieve protein stabilization (Figure 1(B)). Final mutations selection was obtained by merging and ranking the best outcomes of the two methods (Figure 1(C)), which led to the identification of three point mutations located in the variable domain.

Preliminary data on mH6 in vivo toxicity seem to indicate a significant decrease in the mutant toxic profile compared to the wild type cardiotoxic H6. This point out the critical role of specific amino acid positions in determining the toxic behaviour.

Interestingly, the melting temperatures of H6 and mH6 monitored by far-UV circular dichroism are comparable $(T_{\rm m}~{\rm H6}\,{=}\,43.9{-}54.5\,^{\circ}{\rm C}~{\rm and}~T_{\rm m}~{\rm mH6}\,{=}\,53.8\,^{\circ}{\rm C}).$ However, unfolding profiles are not perfectly superposable suggesting different unfolding transitions: mH6 appear to have a more defined transition, which possibly reflect a more compact domain organization than H6 three step unfolding [3]. Limited proteolysis data revealed slower proteolytic kinetics for the mutant, hence reduced fold stability.

The crystal structure of mH6 has been determined at 2.1 Å resolution, displaying a homodimeric organization common to all LC dimers. Secondary structure superposition does not show evident differences between the H6 structure and the mH6 one, giving a root-mean-square deviation (RMSD) value of 0.3 Å for the overall dimers superposition, indicating that the mutagenesis has no impact on the native protein organization and dimerization.

Discussion and conclusions

Previous biophysical findings indicate that low fold stability and high protein dynamics correlate with LC amyloidogenicity [3]. Our data showed that the toxicity of a soluble LC could be tuned by introducing specific point mutations. This evidence points out the importance of the primary sequence for cardiotoxic LCs. Moreover, our data strengthen the hypothesis that links LCs toxic behaviour in vivo to thermal stability and dynamics.

Disclosure statement

No potential conflict of interest was reported by the authors.

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ORCID

Pietro Sormanni http://orcid.org/0000-0002-6228-2221

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