Expression of enzymatically inactive wasp venom phospholipase A1 in Pichia pastoris

Borodina, Irina; Jensen, Bettina M.; Wagner, Tim; Abou Hachem, Maher; Søndergaard, Ib; Poulsen, Lars Kærgaard

Publication date: 2011

Citation (APA):
Expression of enzymatically inactive wasp venom phospholipase A1 in *Pichia pastoris*

Irina Borodina*, Bettina M. Jensen1, Tim Wagner1, Maher A. Hachem2, Ib Søndergaard2 and Lars K. Poulsen2

(1) Center for Microbial Biotechnology, Institute of Systems Biology, Technical University of Denmark, Soeltofts plads 223, 2800 Kgs. Lyngby, Denmark

(2) Allergy Clinic, Dermato-Allergological Dept. K, CUH-Gentofte, Rigshospitalet Dept 7551, Blegdamsvej 9, 2100 København Ø, Denmark

(3) Enzyme and Protein Center, Institute of Systems Biology, Technical University of Denmark, Soeltofts plads 224, 2800 Kgs. Lyngby, Denmark

*ib@bio.dtu.dk

Wasp venom allergy is the most common insect venom allergy in Europe. It is manifested by large local reaction or anaphylactic shock occurring after a wasp sting. The allergy can be treated by specific immunotherapy with whole venom extracts. Wasp venom is difficult and costly to obtain and is a subject to composition variation, therefore it can be advantageous to substitute it with a cocktail of recombinant allergens. One of the major venom allergens is phospholipase A1, which so far has been expressed in *Escherichia coli* and in insect cells. Our aim was to produce the protein in secreted form in yeast *Pichia pastoris*, which can give high yields of correctly folded protein on defined minimal medium and secretes relatively few native proteins simplifying purification.

Residual amounts of enzymatically active phospholipase A1 could be expressed, but the venom protein had a deleterious effect on growth of the yeast cells. To overcome the problem we introduced three different point mutations at the critical points of the active site, where serine137, aspartate165 or histidine229 were replaced by alanine (S137A, D165A and H229A). All the three mutated forms could be expressed in *P. pastoris*. The H229A mutant did not have any detectable phospholipase A1 activity and was secreted up to the level of 4 mg/L in shake flask culture. It was purified by nickel-affinity chromatography and its identity was confirmed by MALDI-TOF mass spectrometry. The protein could bind IgE antibodies from wasp venom allergic patients and could inhibit the binding of wasp venom to IgE antibodies specific for phospholipase A1 as shown by Enzyme Allergo-Sorbet Test (EAST). Moreover, the recombinant protein was allergenic in a biological assay as demonstrated by its capability to induce histamine release of wasp venom-sensitive basophils.

The recombinant phospholipase A1 presents a good candidate for wasp venom immunotherapy.