

Functional Characterization of the *Pseudomonas aeruginosa* ExoY Virulence Factor

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Purpose of the Experiment *P. aeruginosa* is responsible for the second most common hospital infection. Its ExoY toxin is a promiscuous cyclase similar to other toxins like the *B. anthracis* Edema Factor and the *B. pertussis* CyaA. Functional domains of ExoY are poorly understood. I used forward genetic screening to identify functional residues of ExoY, considering the fact that the enzyme is active when expressed in the yeast *S. pombe* and human cells, while inactive in *E. coli*. **Procedures Used** Using PCR I created a population of mutant alleles of *exoY* which were introduced into an expression vector by gap repair transformation. Screening was done using light microscopy and iodine staining to detect hypomorphic ExoY enzymes. The candidates were sequenced and the single amino acid changes were mapped to a 3D model. The relationship of calmodulin and ExoY was studied by attempting to rescue mutant ExoY activity with mutant alleles of the *S. pombe* calmodulin gene. **Observation/Data/Results** 45 candidates displayed low ExoY activity and were characterized. Most lacked an intact *exoY* gene. From 4 candidates expressing a full-length *exoY* gene, 2 novel single missense mutants were identified; one (R68G), at the active site of the enzyme, the other (Q134R) required for multimerization. I was unable to conclude about the relationship of ExoY and calmodulin, but the identification of 2 distinct co-factor-binding domains on ExoY may explain that calmodulin is necessary, but not sufficient, for ExoY activation. **Conclusions/Applications:** The study can be the basis for development of effective, high affinity ExoY-targeted drugs for treatment of *P. aeruginosa* infections. Future studies will reveal unknown co-factors and promote our knowledge regarding the structure and function.

Awards Won:

Fourth Award of \$500