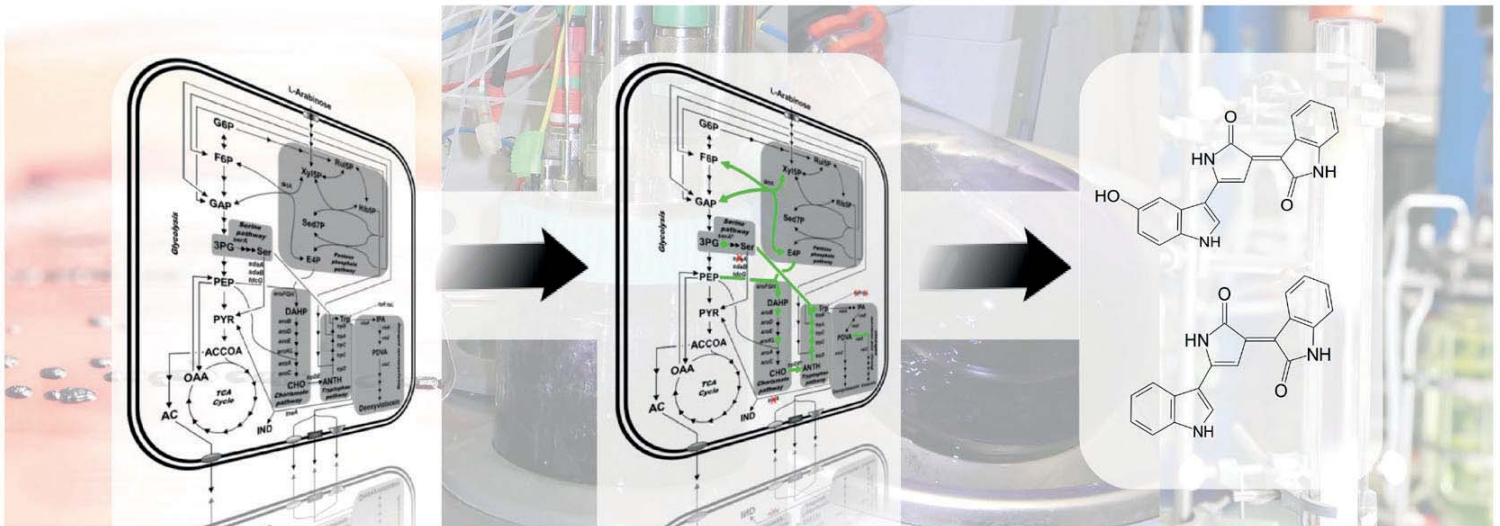




André Luis Rodrigues

Systems metabolic engineering of *Escherichia coli* for production of violacein and deoxyviolacein



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“Reality is merely an illusion, albeit a very persistent one.”

Albert Einstein





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Poblete-Castro, I., Binger, D., Rodrigues, A. L., Becker, J., Dos Santos, V., Wittmann, C., 2013. In-silico-driven metabolic engineering of *Pseudomonas putida* for enhanced production of poly-hydroxyalkanoates. *Metabolic Engineering*. 15, 113-123.

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