

# Producing the Polyketide Antibiotic Actinorhodin From Azo Dye Waste: In silico Design and Optimization of a Synthetic Pathway in *Pseudomonas putida* KT2440

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Recent advances in synthetic biology have enabled the conversion of industrial wastes into high-value chemical products, including specialized metabolites such as polyketides. Azo dyes, a major pollutant commonly discharged by food, textile, and pharmaceutical industries, have the potential to be repurposed as a feedstock for biomanufacturing. Here, I sought to design a synthetic pathway in *Pseudomonas putida* KT2440 capable of recycling azo compounds into the polyketide actinorhodin (ACT). The synthetic pathway was constructed by combining a set of reactions identified from publicly-available databases. Flux balance analysis was used to test the metabolic potential and limitations of the synthetic pathway in *P. putida* using a modified version of the iJN1463 genome-scale metabolic model. The algorithms Optknock, Optgene, and FSEOF were employed to examine whether coupling dye consumption and ACT production with growth could be a suitable optimization strategy. It was revealed that while dye degradation is always coupled with growth, growth-coupling for ACT production is theoretically inefficient. Thus, a synthetic genetic circuit was constructed to decouple growth and ACT production in a two-stage fermentation scenario. Five refactored operons were designed to modularize the ACT gene cluster and avoid overlapping of native and synthetic regulatory elements. To express the system in *P. putida*, the regulatory circuit and refactored operons were integrated into pRO1600 *Pseudomonas*-specific and pBBR1 broad-host-range plasmids using Gibson Assembly. This research demonstrates that the conversion of azo dyes into actinorhodin in *P. putida* is feasible. Moreover, this work establishes a systematic workflow and testable genetic designs to be further investigated experimentally.