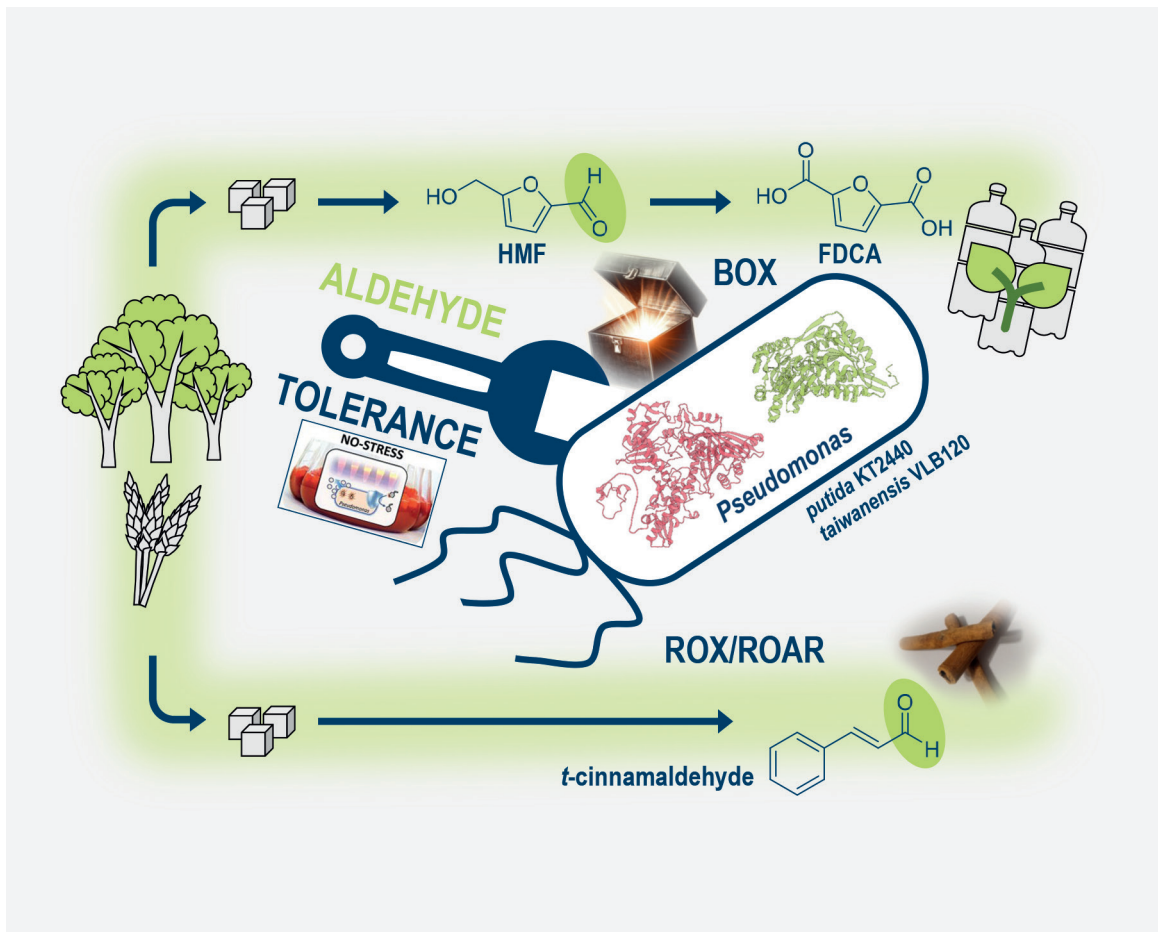




Tolerance engineering of *Pseudomonas* for the efficient conversion and production of aldehydes

Thorsten Lechtenberg

Schlüsseltechnologien / Key Technologies

Band / Volume 292

ISBN 978-3-95806-817-9

Forschungszentrum Jülich GmbH
Institut für Bio- und Geowissenschaften (IBG)
Biotechnologie (IBG-1)

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Schriften des Forschungszentrums Jülich
Reihe Schlüsseltechnologien / Key Technologies

Band / Volume 292

ISSN 1866-1807

ISBN 978-3-95806-817-9

Contents

Publications	I
List of abbreviations	III
List of figures	VI
List of tables	IX
Summary	XIV
Zusammenfassung	XV
1. Introduction	1
1.1. Biocatalysis – Exploring nature’s chemical toolbox.....	1
1.2. Tolerance – A multifaceted prerequisite for efficient and competitive biotechnological processes	3
1.3. Aldehydes	4
1.3.1. Aldehydes in biology	7
1.3.1.1. Aldehyde toxicity.....	8
1.3.2. HMF and furfural – Green chemistry intermediates	9
1.3.3. FDCA as a plant-sourced polymer building block for PEF – Achieving fully bio-based plastics	12
1.3.3.1. Industrial production	13
1.3.3.2. Microbial catalysis as a sustainable alternative.....	14
1.3.4. Microbial degradation of furanic aldehydes and their derivatives	16
1.3.4.1. Genetic basis of the metabolic pathway.....	17
1.3.4.2. The degradation pathway and its biochemistry	19
1.4. <i>Pseudomonas</i> – A multi-talented and robust cell factory	24
1.5. Aim, scope, and outline of this thesis	26

2. Publications and manuscripts	29
2.1. Engineering 5-hydroxymethylfurfural (HMF) oxidation in <i>Pseudomonas</i> boosts tolerance and accelerates 2,5-furandicarboxylic acid (FDCA) production	31
2.1.1. Abstract.....	32
2.1.2. Introduction	34
2.1.3. Results and discussion	36
2.1.3.1. Identification of HMF-oxidizing enzymes in <i>P. taiwanensis</i> VLB120 and <i>P. putida</i> KT2440	36
2.1.3.2. Identification of HMF-reducing enzymes in <i>P. taiwanensis</i> VLB120	39
2.1.3.3. PaoEFG and AldB-I are crucial for tolerance towards HMF	41
2.1.3.4. Tolerance engineering enables GRC1 to withstand higher HMF concentrations	42
2.1.3.5. Optimized FDCA production	45
2.1.4. Conclusion and outlook.....	49
2.1.5. Materials and methods.....	50
2.1.5.1. Strains and culture conditions.....	50
2.1.5.2. Plasmid cloning and strain engineering	51
2.1.5.3. Analytical methods	52
2.1.5.4. Whole-cell HMF conversion assays in shake flasks.....	52
2.1.5.5. Whole-cell HMF conversion assays in 24-deepwell microplates (Duetz-System)	53
2.2. Life with toxic substrates – PaoEFG forms the basis for growth of <i>Pseudomonas taiwanensis</i> VLB120 and <i>Pseudomonas putida</i> KT2440 on aromatic aldehydes	55
2.2.1. Abstract.....	56
2.2.2. Introduction	57
2.2.3. Results and discussion	59
2.2.3.1. PaoEFG is essential for growth on 4-hydroxybenzaldehyde and vanillin.....	59
2.2.3.2. Genomic integration of the <i>hmf</i> -cluster into <i>P. taiwanensis</i> VLB120 GRC1 and <i>P. putida</i> KT2440 enabled growth on furfural, FA, and FDCA, but not on HMF and HMFA	61

2.2.3.3.	Promoting growth on HMF and HMFA by adaptive laboratory evolution (ALE)	64
2.2.3.4.	Whole-genome resequencing of the evolved strains.....	66
2.2.3.5.	Evolved and tolerance-engineered GRC1 PVLB_23545-40::P _{tac} _hmfFGABCDEHT (TL_666 BOX-C/P) strains display superior growth on HMF compared to the native degrader <i>Paraburkholderia caribensis</i>	69
2.2.4.	Conclusion	72
2.2.5.	Materials and methods	72
2.2.5.1.	Strains and culture conditions.....	72
2.2.5.2.	Whole genome sequencing	73
2.2.5.3.	Plasmid cloning and strain engineering	74
2.3.	Improving 5-(hydroxymethyl)furfural (HMF) tolerance of <i>Pseudomonas taiwanensis</i> VLB120 by automated adaptive laboratory evolution (ALE)	76
2.3.1.	Abstract.....	77
2.3.2.	Introduction	78
2.3.3.	Results and discussion.....	80
2.3.3.1.	Increasing HMF tolerance of oxidation-deficient GRC1 ROX by ALE	80
2.3.3.2.	Characterization of evolved strains.....	82
2.3.3.3.	Reverse engineering	84
2.3.3.4.	The benefit of <i>mexT</i> deletion lies in preventing <i>mexEF-oprN</i> expression.....	87
2.3.4.	Conclusion	89
2.3.5.	Materials and methods	90
2.3.5.1.	Strains and culture conditions.....	90
2.3.5.2.	Robotics-assisted ALE.....	91
2.3.5.3.	Whole-genome sequencing	91
2.3.5.4.	Plasmid cloning and strain engineering	92
2.3.5.5.	Analytical methods	92
2.4.	<i>De novo</i> production of <i>t</i> -cinnamaldehyde by engineered solvent-tolerant and oxidation-deficient <i>Pseudomonas taiwanensis</i> VLB120	94
2.4.1.	Abstract.....	95
2.4.2.	Introduction	96

2.4.3. Results and discussion	96
2.4.4. Conclusion	100
2.4.5. Materials and methods.....	100
2.4.5.1. Media and culture conditions	100
2.4.5.2. Plasmid construction and genomic modifications.....	101
2.4.5.3. Analytics.....	101
3. General discussion and perspectives	103
3.1. Whole-cell biocatalytic FDCA production.....	103
3.2. Aromatic aldehydes as products or intermediates of microbial cell factories	108
3.3. Concluding remarks	115
4. References	117
5. Appendix	136
5.1. Supplementary material to chapter 2.1.....	136
5.2. Supplementary material to chapter 2.2.....	146
5.3. Supplementary material to chapter 2.3.....	148
5.4. Supplementary material to chapter 2.4.....	158
Table S1: Strains used in this thesis.....	159
Table S2: Plasmids used in this thesis.	167
Table S3: Oligonucleotides (name, sequence, and description)	174
Danksagung	183
Eidesstattliche Erklärung	185

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