Biotransformations from and to methylated flavonoids

Subtitle

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noch nicht bekannt

Dissertation

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2	BWEIGEL: Überprüfen!	1
3	BWEIGEL: To introduction	2
4	BWEIGEL: Lysozym struktur	2
5	BWEIGEL: nochmal auseinanderklamüsern wegen den konzentrationen und eingesetzten enzymmengen	2
7	BWEIGEL: complete	2
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Preface

1 Abstracts

21.1 English Abstract

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malesuada fames ac turpis egestas. Mauris ut leo. Cras viverra metus rhoncus
sem. Nulla et lectus vestibulum urna fringilla ultrices. Phasellus eu tellus sit amet
tortor gravida placerat. Integer sapien est, iaculis in, pretium quis, viverra ac, nunc.
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varius orci eget risus. Duis nibh mi, congue eu, accumsan eleifend, sagittis quis,
diam. Duis eget orci sit amet orci dignissim rutrum.

14 1.2 Deutsche Zusammenfassung

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Thesis

2 Introduction

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S ome introductionary text
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2.1 Natural products and secondary metabolites

- 42.1.1 General
- **5 2.1.2 Classes of natural products**
- ⁶ Terpenoids and Steroids
- 7 ... here is some text
- 8 Polyketides and non-ribosomal peptides
- 9 ... here is some text
- 10 Alkaloids
- 11 ... here is some text
- 12 Phenylpropanoids
- 13 ... here is some text

14 2.2 Alkylating reactions in nature

- 15 2.2.1 Methylation
- 16 2.2.2 Prenylation

- 12.2.3 Glycosylation
- 22.3 Usage and expansion of natures reaction tool-
- box
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- **5 2.3.2 Methyl transferases**
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- Esterases/Lipases
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- 13 2.4 Conclusion

14 **C**

3 Material And Methods

 $_2$ Within this section percentages refer to volume per volume (v/v) percentages unless $_3$ otherwise specified.

43.1 Materials

53.1.1 Chemicals

Enzymes and buffers used for molecular cloning were obtained from Life Technologies (Darmstadt, Germany), unless otherwise noted. Flavonoid HPLC standards were purchased from Extrasynthese (Genay, France). Deuterated solvents were aquired from Deutero GmbH (Kastellaun, Germany). Solvents, purchased from VWR (Poole, England), were distilled in-house before use.

All other chemicals were obtained from either Sigma-Aldrich (Steinheim, Germany), Applichem (Darmstadt, Germany), Carl Roth (Karlsruhe, Germany) or Merck (Darmstadt, Germany).

14 3.1.2 Commonly used solutions and buffers

 50×5052 25 % glycerol, 2.5 % (w/v) glucose, 10 % (w/v) α -lactose binding buffer 50 mM Tris/HCl, 500 mM NaCl, 10 % glycerol, 2.5 mM imidazole

pH 7

elution buffer 50 mM Tris/HCl, 500 mM NaCl, 10 % glycerol, 250 mM imida-

zole pH 7

lysis buffer $50\,\mathrm{mM}$ Tris/HCl, $500\,\mathrm{mM}$ NaCl, $10\,\%$ glycerol, $2.5\,\mathrm{mM}$ imidazole,

0.2 % Tween-20 pH 7

 $1\,\text{M MMT pH 4 (10\times)} \qquad 26.8\,\text{g/l L-malic acid, } 78.1\,\text{g/l MES, } 26.8\,\text{g/l Tris, } 2.1\,\%\,\,10\,\text{M HCl} \\ 1\,\text{M MMT pH 9 (10\times)} \qquad 26.8\,\text{g/l L-malic acid, } 78.1\,\text{g/l MES, } 26.8\,\text{g/l Tris, } 6.7\,\%\,\,10\,\text{M}$

NaOH

 $20 \times NPS$ 1 M Na₂HPO₄, 1 M KH₂PO₄, 0.5 M (NH₄)₂SO₄

1 M SSG pH 4 (10×)	14.8 g/l succinic acid, 60.4 g/l $\mathrm{NaH_2PO_4}\cdot\mathrm{H_2O},$ 32.8 g/l glycine, 0.4 % 10 M NaOH
1 M SSG pH 10 (10×)	14.8 g/l succinic acid, 60.4 g/l $\mathrm{NaH_2PO_4}\cdot\mathrm{H_2O},$ 32.8 g/l glycine, 10.3 % 10 M NaOH
5× SDS sample buffer	10% (w/v) SDS, 10 mM β -mercaptoethanol, 20% glycerol, $0.2M$ Tris/HCl pH 6.8, 0.05% (w/v) bromophenolblue
1000× trace elements	50 mM FeCl $_3$, 20 mM CaCl $_2$, 10 mM MnCl $_2$, 10 mM ZnSO $_4$, 2 mM CoCl $_2$, 2 mM CuCl $_2$, 2 mM NiCl $_2$, 2 mM Na $_2$ MoO $_4$, 2 mM Na $_2$ SeO $_3$, 2 mM H $_3$ BO $_3$

Preparation of natural deep eutectic solvent (NADES)

 $_2$ NADES were prepared by adding each component in a round-bottom flask with a stirrer and stirring the mixture at 50 $^{\circ}\text{C}$ with intermittent sonication treatments $_4$ until a clear solution was obtained.

Table 3.1.: NADES-mixtures used within this work.

name	composition	mole ratio	mass fraction (w/w)
PCH	propane-1,2-diol	1:1:1	0.326
	choline chloride water		0.597 0.077
GCH	D-glucose choline chloride	2:5:5	0.314 0.608
	water		0.078

5 3.1.3 Culture media used to grow bacteria

LB-medium	10 g/l NaCl, 10 g/l tryptone, 5 g/l yeast extract, pH 7.5
LB-agar	LB + 1.5% (w/v) agar-agar
TB-medium	$12\mathrm{g/l}$ tryptone, $24\mathrm{g/l}$ yeaxst extract, 0.4% glycerol, $72\mathrm{mM}$
	$\mathrm{K_2HPO_4}$, 17 mM $\mathrm{KH_2PO_4}$
ZY	10 g/l tryptone, 5 g/l yeast extract
ZYP-5052	volume fraction (v/v): 0.928 ZY, 0.05 $20 \times$ NPS, 0.02 $50 \times$ 5052,
	$0.002~1~\mathrm{M~MgSO_4},0.0002~1000\times$ trace elements

63.1.4 Bacterial strains

1 E.coli

BL21(DE3) $F^- ompT \ hsdSB(r_B^-, m_B^-) \ gal \ dcm \ \lambda(DE3)$

Invitrogen, Karslruhe (Germany)

C41(DE3) $F^- ompT \ hsdSB(r_B^-, m_B^-) \ gal \ dcm \ \lambda(DE3)$

Lucigen, Wisconsin (USA)

C43(DE3) $F^- \ ompT \ hsdSB(r_B^-, m_B^-) \ gal \ dcm \ \lambda(DE3)$

Lucigen, Wisconsin (USA)

DH5 α F⁻ Φ 80 $lacZ\Delta$ M15 $\Delta(lacZYA-argF)$ U169 recA1 endA1

 $hsdR17(r_K^-m_K^+) phoA supE44 \lambda^- thi-1 gyrA96 relA1$

Invitrogen, Karlsruhe (Germany)

JM110 rpsL thr leu thi lacY galK galT ara tonA tsx dam dcm

 $glnV44 \Delta(lac-proAB)$ e14- [F' $traD36 proAB^+ lacI^q lacZ\Delta M15$]

 $hsdR17(\mathbf{r}_K^-\mathbf{m}_K^+)$

Martin-Luther-University Halle-Wittenberg

JW1593 $rrnB \Delta lacZ4787 \ HsdR514 \Delta (araBAD)$ 568 rph-1 $\Delta ydgG \ (Kan^R)$ (BW25113 derivative) Keio Collection, National Institute of Genetics (Japan)

MG1655 $F^- \lambda^- ilvG^- rfb-50 rph-1$

DSMZ, Hamburg (Germany)

One Shot TOP10 $F^- \Phi 80 lac Z \Delta M15 \Delta (mrr-hsdRMS-mcrBC) recA1 endA1 mcrA$

 $\Delta lacX74 \ araD139 \ \Delta (ara-leu)7697 \ galU \ galK \ rpsL \ (Str^R) \ \lambda^- \ nupG$

Invitrogen, Karlsruhe (Germany)

Origami(DE3) $\Delta (ara-leu)$ 7697 $\Delta lacX$ 74 $\Delta phoA$ Pvull phoR araD139 ahpC galE

galK rpsL F'[lac + lacI q pro] (DE3)gor522::Tn10 trxB (Kan^R,

 Str^R , Tet^R)

Novagen, Wisconsin (USA)

Rosetta(DE3) $F^- ompT \ hsdSB(r_B^-, m_B^-) \ gal \ dcm \ \lambda(DE3) \ pRARE \ (Cam^R)$

Novagen, Wisconsin (USA)

Rosetta(DE3) pLysS F^- ompT $hsdSB(r_B^-, m_B^-)$ gal $dcm \lambda(DE3)$ pLysSRARE (Cam^R)

Novagen, Wisconsin (USA)

T7 Express fhuA2 lacZ::T7 gene1 [lon] ompT gal sulA11 R(mcr-

73:: $miniTn10-Tet^S$)2 [dcm] $R(zgb-210::Tn10-Tet^S$) endA1

Δ(*mcrC-mrr*)114::*IS*10 NEB, Massachusetts (USA)

² Agrobacterium tumefaciens

GV3101 chromosomal background: C58, marker gene: *rif*, Ti-plasmid:

cured, opine: nopaline Sylvestre Marillonet, IPB

3.1.5 Plasmids

Table 3.3.: Plasmids used in this work.

name	supplier/source
pACYCDuet-1	Merck, Darmstadt (Germany)
pCDFDuet-1	Merck, Darmstadt (Germany)
pET-20b(+)	Merck, Darmstadt (Germany)
pET-28a(+)	Merck, Darmstadt (Germany)
pET-32a(+)	Merck, Darmstadt (Germany)
pET-41a(+)	Merck, Darmstadt (Germany)
pQE30	QIAGEN, Hilden (Germany)
pUC19	Invitrogen, Karslruhe (Germany)

23.1.6 Oligonucleotides and synthetic genes

Oligonucleotides and primers were ordered from Eurofins Genomics (Ebersberg, Germany). The purity grade was *high purity salt free* (HPSF). Synthetic genes or gene fragments were obtained from GeneArt® (Life Technologies, Darmstadt, Germany) or Eurofins Genomics (Ebersberg, Germany).

Table 3.4.: Primers used in this work. Recognition sites for endonucleases are underlined. Positions used for site directed mutagenesis are in lower case font.

name	sequence (5'→3')	cloning site
somt1	TTG AAG ACA AAA TGG CTT CTT CAT TAA ACA ATG GCC G	BpiI
somt2	TTG AAG ACA AGG ACA CCC CAA ATA CTG TGA GAT CTT CC	BpiI
somt3	TTG AAG ACA AGT CCT TAG GAA CAC CTT TCT GGG AC	BpiI
somt4	TT <u>G AAG AC</u> A AAA GCT CAA GGA TAG ATC TCA ATA AGA GAC	BpiI
pfomt1.fw	CAG AGA GGC cTA TGA GAT TGG CTT GC	
pfomt1.rv	GCA AGC CAA TCT CAT AgG CCT CTC TG	
pfomt2.fw	CAT ATG GAT TTT GCT GTG ATG AAG CAG GTC	NdeI
pfomt2.rv	GAA TTC AAT AAA GAC GCC TGC AGA AAG TG	EcoRI
pRha1.fw	CTC TAG CAG ATC TCG GTG AGC ATC ACA TCA CCA CAA TTC	BglII
pRha1.rv	CAA TTG A <u>GG ATC C</u> CC ATT TTA ACC TCC TTA GTG	BamHI
pUC1.fw	GCG TAT TGG Gag aTC TTC CGC TTC CTC	
pUC1.rv	GAG GAA GCG GAA GAt ctC CCA ATA CGC	

3.1.7 Instruments

CD-spectrometer Jasco J-815 (Eaton, USA)

electrophoresis (horizon- Biometra Compact XS/S (Göttingen, Germany)

tal)

electrophoresis (vertical) Biometra Compact M (Göttingen, Germany)

Biometra Minigel-Twin (Göttingen, Germany)

FPLC ÄKTA purifier (GE Healthcare, Freiburg, Germany)
GC/MS GC-MS-QP2010 Ultra (Shimadzu, Duisburg, Germany)
HPLC VWR-Hitachi LaChrom Elite (VWR, Darmstadt, Germany)

ITC MicroCal iTC200 (Malvern, Worcestershire, UK)

plate-reader SpectraMax M5 (Molecular Devices, Biberach, Germany)

NMR-spectrometer Varian Unity 400 (Agilent, Böblingen, Germany)

Varian VNMRS 600 (Agilent, Böblingen, Germany)

photospectrometer Eppendorf Biophotometer Plus (Hamburg, Germany)

JASCO V-560 (Eaton, USA)

Colibri Microvolume Spectrometer (Biozym, Hess. Olden-

dorf, Germany)

centrifuges Eppendorf 5424 (Hamburg, Germany)

Hettich Mikro 120 (Kirchlengern, Germany)

Beckman Avanti J-E, Beckman Allegra X-30R (Krefeld,

Germany)

centrifuge rotors Beckman JA-10, JA-16.250, JS-4.3 (Krefeld, Germany)

23.1.8 Software

All mathematical and statistical computations and graphics were done with the R software (versions 3.1.X, http://cran.r-project.org/) [41]. Visualizations of macromolecules were arranged using the PyMol Molecular Graphics System, version 1.7.0.0 (Schrödinger, New York, USA).

Physicochemical calculations and calculations of different molecular descriptors were performed using Marvin Beans 15.4.13.0 (ChemAxon, Budapest, Hungary) and Molecular Operating Environment 2008.10 (Chemical Computing Group, Montreal, Canada). Special software used for X-ray crystal structure solution is discussed

12 3.2 Molecular Biology

separately in the corresponding section (3.5).

¹³ Basic molecular biology methods like polymerase chain reaction (PCR), DNA re-¹⁴ striction/ligation, DNA gel electrophoresis, preparation of competent cells and transformation were performed based on the protocols summarized by Sambrook and Russell [47].

Plasmid DNA was isolated using the QIAprep® Spin Miniprep Kit (QIAGEN, Hilden, Germany) according to the manufacturer's instructions.

⁵ In vitro site-directed mutatgenesis was set-up according to the protocol of the ⁶ QuikChange[™] Site-Directed Mutagenesis kit [2] offered by Agilent Technologies ⁷ (Santa Clara, USA).

Nucleotide fragments obtained by PCR, restriction/ligation procedures or excision from electrophoresis gels were purified and concentrated using the *Nucleospin Gel* and *PCR Clean-up* kit provided by Machery-Nagel (Düren, Germany) according to the instructions provided by the manufacturer.

12 3.2.1 Golden Gate Cloning

The Golden Gate cloning procedure is a one-pot method, meaning the restriction digestion and ligation are carried out in the same reaction vessel at the same time [27, 15]. Consequently PCR-fragments, destination vector, restriction endonuclease and ligase are added together in this reaction. The methodology employs type II restriction enzymes, which together with proper design of the fragments allow for a ligation product lacking the original restriction sites.

 19 For digestion/ligation reactions of fragments containing BpiI sites, 20 fmol of each 20 fragment or vector, together with 5 U of BpiI and 5 U of T4 ligase were combined in 21 a total volume of 15 μ l 1× ligase buffer. For fragments to be cloned via BsaI sites, 22 BpiI in the above reaction was substituted by 5 U BsaI.

 23 The reaction mixture was placed in a thermocycler and inbcubated at 37 °C for 24 2 min and 16 °C for 5 min. These two first steps were repeated 50 times over. Finally, the temperature was raised to 50 °C (5 min) and 80 °C (10 min) to inactivate the enzymes.

27 3.2.2 Subcloning of genes

²⁸ All subcloning procedures were performed according to section 3.2 and specifically ²⁹ subsection 3.2.1. Specific steps for the subcloning of any genes discussed can be ³⁰ found in the appendix (p.37). The *pfomt* gene was subcloned from the pQE-30 vector kindly provided by Thomas Vogt (Leibniz-Institute of Plant Biochemistry ³² (IPB), Halle, Germany) into the pET-28a(+) vector. The *somt-2* gene was subcloned ³³ from the pQE-30 vector kindly provided by Martin Dippe (IPB, Halle, Germany) ³⁴ into the pET-28-MC vector.

13.2.3 Transformation of electrocompetent *Agrobacterium*2 tumefaciens cells

³ A 50 μl aliquot of electrocompetent *A. tumefaciens* cells was thawed on ice. (50 to 4 100) ng of plasmid were added, the solution was mixed gently and transferred to a 5 pre-cooled electroporation cuvette. After pulsing $(2.5 \text{ kV}, 200 \Omega)$ 1 ml of lysogeny 6 broth (LB)-medium was added, the mixture transferred to a 1.5 ml tube and incubated for (3 to 4) hours at 28 °C. The culture was centrifuged $(10\,000 \times g, 1\,\text{min})$ and 900 μl supernatant were discarded. The pellet was resuspended in the remaining liquid, plated onto LB-agar plates supplemented with $40\,\mu\text{g/ml}$ rifampicin and $10\,50\,\mu\text{g/ml}$ carbencillin and incubated at 28 °C for $(2\,\text{to}\,3)\,\text{days}$.

3.3 Treatment of plant material

12 3.3.1 Infiltration of Nicotiana benthamiana

Before infiltration *N. benthamiana* plants were pruned, such that only leaves to be infiltrated remained with the plant. 5 ml cultures of transformed *A. tumefaciens* in LB-medium (with $40 \,\mu\text{g/ml}$ rifampicin and $50 \,\mu\text{g/ml}$ carbencillin) were grown over night at $28 \,^{\circ}\text{C}$ and $220 \,\text{rpm}$. OD⁶⁰⁰ of the culture was measured and adjusted to $17 \, 0.2$ by dilution with infiltration buffer ($10 \,\text{mM}$ MES/NaOH, $10 \,\text{mM}$ MgSO₄ pH 5.5). When multiple *A. tumefaciens* transformed with different contructs/plasmids were used for infiltration, the cultures were mixed and diluted using infiltration buffer, such that OD⁶⁰⁰ of each culture in the mix was 0.2. The solution was infiltrated into the abaxial side of *N. benthamiana* leaves using a plastic syringe. The leaf material was harvested after 7 days.

23 3.3.2 Plant material harvest

Infiltrated/Infected areas of *N. benthamiana* leaf material were cut out and grouped by plant number, leaf position (top/bottom) and leaf side (right/left). The grouped clippings were weighed, frozen in liquid nitrogen, ground to a powder, freeze-dried and stored at -80 °C.

28 3.3.3 Extraction of flavonoids from N. benthamiana leaves

Two tips of a small spatula of freeze-dried material (\approx 6 mg), were weighed exactly and extracted with 500 μ l 75 % aqueous methanol containing 1 mM ascorbic acid, 0.2 % formic acid and 0.1 mM flavone (internal standard). Therefore the suspension was vortexed for 30 s, rotated on an orbital shaker for 10 min and vortexed again for

 $_{2}$ 30 s. The suspension was centrifuged (20 000 × g, 4 °C, 10 min) and the supernatant $_{2}$ transferrerd to a new tube, to remove the insoluble plant material. The supernatant $_{3}$ was centrifuged again (20 000 × g, 4 °C, 10 min) and the resulting supernatant was $_{4}$ transferred to a HPLC-vial and stored at $_{2}$ 0 °C until analysis.

3.4 Protein biochemistry

⁶ Stock solutions of antibiotics, IPTG or sugars were prepared according to the pET ⁷ System Manual by Novagen [38], unless otherwise noted.

3.4.1 Determination of protein concentration

Protein concentrations were estimated using the absorption of protein solutions at 280 nm, which is mainly dependent on the amino acid composition of the protein studied [19]. Extinction coefficients of proteins were calculated from the amino acid sequence using the ExpPASy servers's ProtParam tool [18].

Table 3.6.: Calculated extinction coefficients of proteins used in this work.

protein/enzyme	$arepsilon_{280\mathrm{nm}}^{1\mathrm{g/l}}$ in $\mathrm{ml}\mathrm{mg}^{-1}\mathrm{cm}^{-1}$
PFOMT (reduced)	0.714
PFOMT Y51K N202W (reduced)	0.852
SOMT-2 (oxidized)	1.263
SOMT-2 (reduced)	1.247
COMT	

3.4.2 Protein production test (expression test)

- The heterologous production of proteins in *E. coli* was assessed in a small scale
- protein production test, henceforth called expression test. Single colonies of *E. coli*
- transformed with the constructs to be studied were used to inoculate a 2 ml starter
- rd culture in LB-medium containing the appropriate antibiotics. The working con-
- centrations of antiobiotics used was as follows: $200\,\mu\text{g/ml}$ ampicillin, $150\,\mu\text{g/ml}$
- 19 kanamycin, 50 μg/ml chloramphenicol, 20 μg/ml tetracycline.
- The starter culture was allowed to grow at 37 °C and 200 rpm over night. A 5 ml
- $_{\mbox{\tiny 21}}$ sampling culture of the medium to be studied containing the appropriate antibiotics
- 22 was prepared. The media tested included LB, terrific broth (TB) and auto-induction
- 23 media like ZYP-5052. The sampling culture was inoculated to an OD^{600} of 0.075
- ²⁴ using the starter culture and incubated at different temperatures and 200 rpm in

a shaking incubator. 1 mM isopropyl-D-thiogalactopyranosid (IPTG) was added when the OD⁶⁰⁰ reached 0.6-0.8, if appropriate for the studied construct. 1 ml samples were removed after different times of incubation (e.g. 4, 8, 12 hours), subfractionated (3.4.3) and analyzed via SDS-polyacrylamide gel electrophoresis (PAGE) (3.4.6).

⁶ Exact specifications of growth conditions (e.g. temperature, time, constructs) are ⁷ discussed in the individual sections.

3.4.3 Protein subfractionation

The protein subfractionation procedure described herein was adapted from the protocol described in the pET Manual [38]. Overall 5 protein subfractions can be obtained, including total cell protein, culture supernatant (medium) protein, periplasmic protein, solube cytoplasmic protein and insoluble protein.

The OD⁶⁰⁰ of the culture sample was measured and the cells harvested by centrifu-

The OD⁶⁰⁰ of the culture sample was measured and the cells harvested by centrifugation at $10\,000\times g$, $4\,^{\circ}$ C for 5 minutes. The protein in the supernatant medium was concentrated by precipitation with trichloro acetic acid (TCA) (3.4.4) for SDS-PAGE analysis. The periplasmic protein was prepared (3.4.5) and also concentrated by TCA precipitation for SDS-PAGE. Cells were lysed by resuspending the cell pellet in (OD⁶⁰⁰ × V × 50) µl of bacterial protein extraction reagent (B-PER) and vortexing vigorously for 30 s. The suspension was incubated at room temperature (RT) for 30 min to assure complete lysis. To separate insoluble protein and cell debris from the soluble cytosolic protein, the suspension was centrifuged at $10\,000\times g$ and $4\,^{\circ}$ C for 10 min. Soluble cytoplasmic protein was contained in the supernatant, whereas insoluble protein remained in the pellet. For SDS-PAGE analysis of the insoluble protein, the pellet was resuspended in the same volume of B-PER. To obtain only the total cell protein fraction, the preparation of periplasmic and soluble cytosolic protein was omitted. Sample volumes of $10\,\mu$ l of each fraction were used for SDS-PAGE analysis.

28 3.4.4 Protein sample concentration by TCA precipitation

Diluted protein samples were concetrated by TCA precipitation in microcentrifuge tubes. Therefore 0.1 volume (V) of 100% (w/v) TCA in water was added to the clarified sample, which was then vortexed for 15 s and placed on ice for a minimum of min. The sample was centrifuged at $14000\times g$, 4° C for 15 min. The supernatant was discarded and the pellet was washed twice with 0.2 V ice-cold acetone. The acetone was removed and the pellet set to air-dry in an open tube. After drying, the protein pellet was resuspended in 0.1 V phosphate buffered saline (PBS) containing $1\times$ SDS-sample buffer by heating to 85 °C and vigorous vortexing, to achieve a

 $_1$ 10 × concentration. After resuspension the sample was analyzed by SDS-PAGE or $_2$ stored at $-20\,^{\circ}$ C until use.

3.4.5 Preparation of periplasmic protein

Target proteins may be directed to the periplasmic space by N-terminal signal sequences like pelB or DsbA/C [31]. The periplasma is, other than the cytosol, an oxidizing environment and often used for the production of proteins containing dilsufide linkages. The preparation of periplasmic protein was accomplished by an osmotic shock protocol modified from Current Protocols in Molecular Biology [5]. The cell pellet was resuspended in the same volume as the culture sample of 30 mM tris-HCl, 20 % (w/v) sucrose, pH 8 and 1 mM ethylenediaminetetraacetic acid (EDTA) was added. The suspension was stirred for 10 min at RT and the cells were collected by centrifugation at $10\,000\times g$, $4\,^{\circ}$ C for 10 min. The supernatant was discarded and the cell pellet was resuspended in the same volume of ice-cold 5 mM MgSO₄. The suspension was stirred for 10 min on ice, while the periplasmic proteins were released into the solution. The cells were collected by centrifugation as before. Periplasmic proteins were contained in the supernatant.

17 3.4.6 Discontinous SDS-polyacrylamide gel electrophoresis (SDS-PAGE)

19 The analysis of samples via SDS-PAGE was realized via the discontinous system 20 first described by Laemmli, which allows separation of proteins based on their 21 electrophoretic mobility, which in turn depends on their size [29]. 22 The SDS-PAGE procedure was carried out according to standard protocols described 23 by Sambrook and Russell [47]. Very dilute and/or samples with high ionic strength 24 were concentrated and/or desalted by the TCA precipitation procedure described in 25 subsection 3.4.4. Generally a 10 % (acrylamide/bisacrylamide) running gel combined 26 with a 4 % stacking gel was used. Reducing SDS-PAGE sample buffer was added to 27 the protein sample to be analyzed, whereafter the sample was heated to 95 °C for 28 5 min, to allow for total unfolding of the protein. After cooling to RT the samples ²⁹ were transferred into the gel pockets for analysis. The PageRuler[™] Prestained 30 Protein Ladder (Life Technologies GmbH, Darmstadt, Germany) was used as a molecular weight (MW) marker and run alongside every analysis as a reference. 32 Gels were stained using a staining solution of 0.25 % Coomassie Brilliant Blue G-33 250 (w/v) in water:methanol:acetic acid (4:5:1) and destained by treatment with 34 water:methanol:acetic acid (6:3:1).

3.4.7 Buffer change of protein samples

The buffer in protein samples was exchanged either by dialysis, or by centrifugal filter concentrators (Amicon[®] Ultra Centrifugal Filter; Merck, Darmstadt, Germany). Large volumes of highly concentrated protein solutions were preferably dialyzed. Respectively, very dilute samples were concentrated and rebuffered using centrifugal concentrators.

⁷ Dialysis was carried out at least twice against a minimum of 100 times the sample ⁸ volume. Dialysis steps were carried out at RT for 2 hours, or over-night at 4 °C. ⁹ Centrifugal concentrators were used according to the manufacturers instructions.

10 3.4.8 Production of recombinant protein

11 Heterologous production of PFOMT

¹² PFOMT was produced as a N-terminally (His)₆-tagged fusion protein. A 2 ml starter ¹³ culture of LB containing 100 μg/ml kanamycin was inoculated with a single colony of E. coli BL21(DE3) transformed with pET28-pfomt and incubated at 37 °C, 220 rpm for 6 hours. The main culture (N-Z-amine, yeast extract, phosphate (ZYP-5052) 16 containing 200 μg/ml kanamycin) was inoculated with the starter culture such that ¹⁷ OD⁶⁰⁰ was 0.05. The culture was incubated in a shaking incubator at 37 °C, 220 rpm ¹⁸ over night (≈16 h). Due to the autoinducing nature of the ZYP-5052 medium, addi-19 tion of IPTG was not neccesary. Cells were harvested by centrifugation at $10\,000 \times g$, ₂₀ 4 °C for 10 min and the supernatant discarded. The pellet was resuspended in 50 mM ²¹ Tris/HCl, 500 mM NaCl, 2.5 mM imidazole, 10 % glycerol pH 7 using a volume of $_{22} \approx 10 \text{ ml/g}$ of cell pellet. The cells were lysed by sonication (70 % amplitude, 1 s on-23 off-cycle) for 30 seconds, which was repeated twice. The crude lysate was clarified ₂₄ by centrifugation at $15\,000 \times g$, $4\,^{\circ}$ C for 15 minutes followed by filtration through a 25 0.45 μm filter. Consequently, the His-tagged PFOMT was purified by immobilized ₂₆ metal affinity chromatography (IMAC) (3.4.10). The eluted PFOMT protein was 27 dialyzed (3.4.7) against 25 mM HEPES, 100 mM NaCl, 5 % glycerol pH 7 and stored 28 at −20 °C until use.

29 Heterologous production of SOMT-2

 $_{30}$ SOMT-2 was produced as a fusion protein with an N-terminal His-tag. A starter $_{11}$ LB-culture (≈ 2 ml) containing 100 µg/ml kanamycin was inoculated with a single colony of *E. coli* BL21(DE3) transformed with pET28MC-somt and incubated at $_{13}$ 37 °C, 220 rpm for 6 hours. The starter culture was used to inoculate the main culture (LB-medium containing 100 µg/ml kanamycin), such that OD $_{10}^{600}$ ≈ 0.05 . The culture was incubated at 37 °C, 220 rpm in a shaking incubator until OD $_{10}^{600}$

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 $_1 \approx 0.6$. Expression was induced by addition of 1 mM IPTG. Incubation continued at $_2$ 37 °C, 220 rpm for 6 hours. Cells were harvested by centrifugation ($10\,000 \times g$, 4 °C, $_3$ 10 min) and used, or stored at $_2$ °C until use. SOMT-2 was produced in inclusion $_4$ bodies (IBs), which were prepared as laid out in subsection 3.4.9.

3.4.9 Preparation of inlusion bodies (IBs)

6 Often, when recombinant protein is produced in high levels in E. coli it is accumulated in so-called inlusion bodies (IBs) [45]. The accumulating IBs consist mainly s of the overproduced target protein, which is inherently quite pure already. IBs can \circ be selectively recovered from *E. coli* cell lysates and can consequently be refolded. 10 IBs were prepared according to a modified protocol by Palmer [39]. The cells were resusupended in 5 ml/g_{cells} IB lysis buffer (100 mM Tris/HCl, 1 mM 12 EDTA pH 7), 0.5 mM phenylmethylsulfonylfluoride (PMSF) was added as protease inhibitor. The solution was homogenized using a tissue grinder homogenizer (Ultra 14 Turrax[®]; IKA[®]-Werke GmbH & Co. KG, Staufen, Germany). 200 μg/ml lysozyme 18 was added to aid in the breakage of cells and the cells were lysed by sonicating thrice at 70 % amplitude (1 s on-off-cycle) for 30 seconds. DNase I (10 µg/ml) was ₁₇ added and the solution was incubated on ice for 10 min. The lysate was clarified by ¹⁸ centrifuging for 1 h at 20 000 \times g, 4 °C. The supernatant was discarded and the pellet $_{19}$ was resuspended in 5 ml/g_{cells} IB wash buffer I (20 mM EDTA, 500 mM NaCl, 2 %₂₀ (w/v) Triton X-100 pH), followed by thorough homogenization. The solution was ₂₁ centrifuged (30 min at 20 000 $\times g$, 4 °C), the supernatant discarded and the pellet 22 was washed twice more. To remove detergent, the pellet was washed twice again 23 with IB washing buffer II (20 mM EDTA, 100 mM Tris/HCl pH 7). The IBs were ²⁴ resuspended in IB solubilization buffer (100 mM Tris/HCl, 5 mM DTT, 6 M GdmCl 25 pH 7), such that the protein concentration was about 25 mg/ml and stored at −20 °C 26 until use.

27 3.4.10 Purification of His-tagged proteins using immobilized metal affinity chromatography (IMAC)

N- or C-terminal oligo-histidine tags (His-tags) are a common tool to ease purification of recombinantly produced proteins. The free electron pairs of the imidazol nitrogens of histidines can complex divalent cations such as Mg²⁺ or Ni²⁺, which are usually immobilized on a matrix of nitrilo triacetic acid (NTA)-derivatives. The affinity of the His-tag is correlated with its length and tagged proteins can simply be eluted by increasing the concentration of competing molecules (e.g. imidazole). His-tagged protein was purified by fast protein liquid chromatography (FPLC) via Ni²⁺- (HisTrap FF crude) or Co²⁺-NTA (HiTrap Talon FF crude) columns, obtained

from GE Healthcare (Freiburg, Germany), following modified suppliers instructions. First the column was equilibrated with 5 column volumes (CV) of binding buffer (50 mM Tris/HCl, 500 mM NaCl, 10 % glycerol, 2.5 mM imidazole pH 7). The sample (generally clarified lysate) was applied to the column using a flow of 0.75 ml/min. Unound protein was removed by washing with 3 CV binding buffer. Unspecifically bound proteins were washed away by increasing the amount of elution buffer (50 mM Tris/HCl, 500 mM NaCl, 10 % glycerol, 250 mM imidazole pH 7) to 10 % (constant for 3 to 5 CV). Highly enriched and purified target protein was eluted with 6 to 10 CV of 100 % elution buffer.

10 3.4.11 Refolding of SOMT-2 on a micro scale using design of experiments (DoE)

- Design of experiments (DoE) and FFD have been successfully used to optimize the refolding conditions of several proteins [55, 3, 6]. Thus, an approach using FFD
- 14 was used to find optimal refolding conditions for SOMT-2.
- 15 Factors studied were pH (buffer), arginine, glycerol, divalent cations, ionic strength,
- 16 redox system, cyclodextrin and co-factor addition. The experimental matrix was
- rg constructed using the FrF2 package (http://cran.r-project.org/web/packages/FrF2/
- 18 index.html) in the R software.

Table 3.7.: Factors used in the construction of the FFD.

factor	symbol		unit	
		-1	+1	
pH	A	5.5	9.5	-
arginine	В	0	0.5	M
glycerol	C	0	10	% (v/v)
divalent cations ¹	D	no	yes	-
ionic strength ²	E	low	high	-
redox state ³	F	reducing	redox-shuffling	-
α -cyclodextrin	G	0	30	mM
SAH	Н	0	0.5	mM

 $^{^{1}}$ no: 1 mM EDTA; yes: 2 mM CaCl $_{2}$, MgCl $_{2}$

²low: 10 mM NaCl, 0.5 mM KCl; high: 250 mM NaCl, 10 mM KCl

³reducing: 5 mM DTT; redox-shuffling: 1 mM glutathione (GSH), 0.2 mM glutathione disulfide (GSSG)

Table 3.8.: Experimental design matrix for the FFD.

Experiment	A	В	С	D	Е	F	G	Н
1	+	+	+	-	-	-	-	+
2	-	-	-	-	-	-	-	-
3	+	-	+	+	-	+	+	-
4	-	+	+	-	+	+	+	-
5	+	+	-	-	+	+	-	-
6	-	+	-	+	+	-	+	+
7	+	+	-	+	-	-	+	-
8	-	-	+	-	+	-	+	+
9	+	-	+	+	+	-	-	-
10	-	-	-	-	-	+	+	+
11	+	-	-	+	+	+	-	+
12	-	+	+	+	-	+	-	+

The buffers were mixed from stock solutions and prepared in 1.5 ml microcentrifuge tubes immediately prior to the experiment. 50 μ l of solubilized SOMT-2 (1 mg/ml) in IB solubilization buffer was added to each buffer followed by a short vortex boost for rapid mixing. The final protein concentration in the refolding reaction was 50 μ g/ml, whereas the remaining GdmCl concentration was \approx 286 mM. The refolding reactions were incubated at RT for 1 hour, followed by an over night incubation at 4 °C. After incubation the refolding reactions were centrifuged (10 000 × g, 4 °C, 10 min) to separate insoluble and soluble protein fractions. The supernatant was transferred to a new tube, whereas the pellet was washed twice with 200 μ l acetone and once with 400 μ l methanol/acetone (1:1). The pellet was resuspended in 100 μ l PBS with 20 μ l SDS-PAGE sample buffer and 10 μ l were used for SDS-PAGE analysis.

100 μl of the supernatant were concentrated using TCA precipitation (3.4.4) and analyzed by SDS-PAGE. The remaining supernatant was rebuffered into 50 mM 1.5 2-[Bis(2-hydroxyethyl)amino]-2-(hydroxymethyl)propane-1,3-diol (BisTris) pH 7.5 1.6 using Amicon® Ultra 0.5 ml centrifugal filters (Merck, Darmstadt, Germany) according to the manufacturers instructions. The pre-weighed collection tubes were re-weighed after recovery and the volume of recovered liquid calculated 1.9 ($\rho \approx 1\,\mathrm{g/cm^3}$). The sample was filled up to 100 μl using 50 mM BisTris pH 7.5 and 1.0 the protein concentration was assessed using the Roti®-Quant protein quantification solution (Carl Roth, Karlsruhe, Germany) according to the manufacturers description. 50 μl of each refolded sample was used for an activity test using 1.2 naringenin as substrate (3.6.3). The reactions were incubated over night and 1.4 stopped by the extraction method. However, before the actual extraction 1 μl of

anthracene-9-carboxylic acid (AC-9) was added as internal standard. The samples were analyzed by high-performance liquid chromatography (HPLC).

4 Assessment of refolding performance

⁵ The performance of each buffer on the refolding of SOMT-2 was examined by ⁶ comparing the SDS-PAGE results, as well as the amount of soluble protein and the ⁷ conversion of substrate. Main effects were analyzed qualitatively using main effects ⁸ plots [7].

3.4.12 Enzymatic production of SAM and SAE

- ¹⁰ S-adenosyl-L-methionine (SAM) and S-adenosyl-L-ethionine (SAE) were prepared according to the method described by Dippe, et. al [13].
- ¹² Preparative reactions (20 ml) were performed in 0.1 M Tris/HCl, 20 mM MgCl₂,
- 200 mM KCl pH 8.0 and contained 7.5 mM adenosine triphosphate (ATP), 10 mM
- ¹⁴ D,L-methionine or D,L-ethionine, for the production of SAM or SAE respectively,
- and 0.2 U S-adenosylmethionine synthase (SAMS) variant I317V. The reaction was
- 16 stopped by lowering the pH to 4 using 10 M acetic acid after 18 h of incubation
- at 30 °C, 60 rpm. After 10 min incubation on ice the solution was centrifuged
- $_{18}$ (15 000 imes g, 10 min) to remove insoluble matter. The supernatant was transferred to
- 19 a round bottom flask, frozen in liquid nitrogen and lyophilized.
- $_{20}$ Crude products were extracted from the pellet using 73 % ethanol and purified
- ²¹ using ion exchange chromatography (IEX). IEX was performed on a sulfopropyl
- 22 sepharose matrix (25 ml) via isocratic elution (500 mM HCl). Before injection, the
- 23 crude extract was acidified to 0.5 M HCl using concentrated hydrochloric acid.
- After elution, the product containing fractions were dried via lyophilization.
- The amount of product was determined by UV/VIS-spectroscopy at 260 nm using the published extinction coefficient of SAM ($\varepsilon_0 = 15\,400\,\mathrm{M}^{-1}\,\mathrm{cm}^{-1}$) after resuspension
- 27 in water [49].

28 3.5 Crystallographic Procedures

29 3.5.1 Crystallization of proteins

- 30 Commercially available crystallization screens were used to find initial crystalliza-
- 31 tion conditions. The tested screens included kits available from Hampton Research
- 32 (Aliso Viejo, USA) and Jena Bioscience (Jena, Germany). Crystallization screens
- were processed in 96-well micro-titer plate (MTP)s, where each well possessed

4 subwells aligned in a 2×2 matrix. The subwells were divided into 3 shallow wells for sitting drop vapour diffusion experimental setups and a fourth subwell, which was deep enough to act as buffer reservoir. This way the performance of each crystallization buffer could be assessed using three different protein solutions with varying concentrations, effectors etc. A pipetting robot (Cartesian Microsys, Zinsser-Analystik; Frankfurt, Germany) was used to mix 200 nl of each, protein and buffer solution, for a final volume of 400 nl. The crystallization preparations were incubated at 16 °C and the progress of the experiment was documented by an automated imaging-system (Desktop Minstrel UV, Rigaku Europe, Kent, UK). Furthermore, fine screens (e.g. for refinement of crystallization conditions) were set up by hand in 24-well MTPs using the hanging drop vapour diffusion method.

12 **PFOMT**

- PFOMT protein was concentrated to (6 to 8) mg/ml and rebuffered to 10 mM
- Tris/HCl pH 7.5 using Amicon® Ultracel centrifugal concentrators (10 kDa MWCO).
- The concentrated protein solution was centrifuged at $14\,000 \times g$, 4°C for 10 min to
- ¹⁶ remove any insoluble material or aggregates. Crystallization screens were set up as described above.
- 18 apo-PFOMT was crystallized using the following conditions 2 M (NH₄) $_2$ SO $_4$,
- $_{19}$ 20 %glycerol. The protein solution contained 0.25 mM SAE, 0.25 mM MgCl $_{2}$,
- $_{20}$ 0.25 mM eriodictyol and 7.53 mg/ml (0.262 mM) PFOMT.

21 Crystallization of proteins using NADES

²² NADES have the potential to be excellent solvents for hydrophobic compounds ²³ such as flavonoids or cinnamic acids [10] and in addition they are able to stabilize ²⁴ and activate enzymes [20].

Four different model proteins (bovine trypsin, hen-egg white lysozyme, proteinase K and *Candida cylindrica* lipase B) were used to assess the capability of NADES for protein crystallization. PCH was tested in a full factorial grid layout using PCH concentrations of (20, 30, 40 and 50) % combined with buffers of different pH. The buffers included 0.1 M sodium acetate pH (4.5 and 5.5), 0.1 M sodium citrate pH 6.5, 0.1 M 2-[4-(2-hydroxyethyl)piperazin-1-yl]ethanesulfonic acid (HEPES)/NaOH pH (7 and 7.5) and 0.1 M Tris/HCl pH 8.5. Thus, the full factorial design had a size of $4 \times 6 = 24$ different conditions. Protein solutions were prepared from lyophilized protein and were as follows: 90 mg/ml trypsin in 10 mg/ml benzamidine, 3 mM CaCl₂; 75 mg/ml lysozyme in 0.1 M sodium acetate pH 4.6; 24 mg/ml proteinase K in 25 mM Tris/HCl pH 7.5 and 6 mg/ml lipase B in water. For crystallization 2 μ l enzyme solution and 1 μ l reservoir buffer were mixed

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3.5. Crystallographic Procedures

and set up in a hanging drop experiment on a 24-well MTP. The experiments were set up at 4 °C.

3.5.2 Data collection and processing

⁴ Crystallographic data were collected at the beamline of the group of Professor Stubbs (MLU, Halle, Germany). The beamline was equipped with a rotating anode 6 X-ray source MicroMax007 (Rigaku/MSC, Tokio, Japan), which had a maximum 7 power of 0.8 kW (40 kV, 20 mA) and supplied monochromatic Cu-K_α-radiation with 8 a wavelength of 1.5418 Å. Diffraction patterns were detected with a Saturn 944+ 9 detector (CCD++, Rigaku/MSC, Tokio, Japan).

Indexing and integration of the reflexes via Fourier transformation (FT) was accomplished using *XDS* [25, 24, 26] or *MOSFLM* [40]. *Scala* [16], which is integrated in the Collaborative Computational Project No. 4 (CCP4)-Suite, was used for scaling of the intensities.

14 3.5.3 Structure solution

For the determination of the electron density $\rho(\mathbf{r})$, where \mathbf{r} is the positional vector, from the diffraction images by FT two terms are neccessary as coefficients; the structure factor amplitudes, $F_{\text{obs}}(\mathbf{h})$ and the phase angles or phases, $\alpha(\mathbf{h})$, where \mathbf{h} is the reciprocal index vector. The structure factor amplitudes can be directly determined from the measured and corrected diffraction intensities of each spot. However, the phase information is lost during the detection of the diffracted photons and there is no direct way to determine the phases. This constitutes the so-called phase problem. Thus, additional phasing experiments are necessary in order to obtain the phases. A variety of phasing experiments are available, which include marker atom substructure methods, density modification and molecular replacement (MR) techniques [46]. Phases of the structures herein were exclusively determined by MR [43, 44].

²⁷ MR was performed using the software *Phaser* [32, 33], which is included in the ²⁸ CCP4-Suite [56]. A previously published PFOMT structure (PDB-code: 3C3Y [28]) ²⁹ was used as a template during MR procedure for the PFOMT structure solution.

For the MR of the lysozyme structure the PDB-entry 4NHI was used.

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31 3.5.4 Model building, refinement and validation

³² Macromolecular model building and manipulation, as well as real space refine-³³ ment and Ramachandran idealization were performed using the Crystallographic Object-Oriented Toolkit (*Coot*) software [14]. Structure refinement was done using the software REFMAC5 [36, 53] as part of the CCP4-suite or the Phyton-based Hierarchial Environment for Integrated Xtallography (PHENIX) [1]. Validation of the structures was carried out using the web service MolProbity (*http://molprobity.biochem.duke.edu/*) [8]. Structure visualization and the preparation of figures was performed using PyMOL (Schrödinger, New York, USA).

73.5.5 In silico substrate docking

In silico molecular docking studies were performed using the AutoDock Vina 1.1.2 or AutoDock 4.2.6 software in combination with the AutoDockTools-Suite (http://autodock.scripps.edu/) [21, 35, 52]. Substrates were docked into the PFOMT structure with the PDB-code 3C3Y. The grid box, which determines the search space, was manually assigned to center at 1.581, 5.196 and 25.718 (x, y, z) and had size of (22, 20 and 25) Å (x, y, z). The exhaustiveness of the global search for AutoDock Vina was set to 25, whereas the rest of the input parameters were kept at their defaults.

16 3.6 Analytics

17 3.6.1 Recording of growth curves

Starter cultures (\approx 2 ml) of the transformed *E. coli* cells were prepared in the medium to be studied, containing the appropriate antibiotics. The cultures were incubated at 37 °C, 200 rpm over night and harvested by centrifugation ($5000 \times g$, 4 °C, 5 min). The pellet was resuspended in 15 ml PBS and the suspension centrifuged ($5000 \times g$, 22 4 °C, 5 min). The supernatant was discarded and the washing step repeated once more. The washed pellet was resuspended in 2 ml of the medium to be studied with the appropriate antibiotics and the OD⁶⁰⁰ was measured. Three independent 50 ml cultures of the medium containing the appropriate antibiotics were inoculated such that OD⁶⁰⁰ \approx 0.05 using the washed cell suspension. The cultures were incubated at the conditions to be studied and sampled at appropriate intervals of time (\approx 1 h). One ml samples were kept on ice until all samples were aquired. 100 µl aliquots of the samples were transferred into a clear MTP and the OD⁶⁰⁰ was measured. Green fluorescent protein (GFP) fluorescence was measured accordingly, but the MTP used was opaque. Exitation (λ ^{ex}) and emission (λ ^{em}) wavelengths were (470 and 510) nm respectively.

3.6.2 In vitro determination of glucose

The glucose concentration in clarified, aqueous samples was determined by a modified version of the glucose assay kit procedure provided by Sigma-Aldrich [50]. Glucose oxidase (GOD) oxidizes D-glucose to gluconic acid, whereby hydrogen peroxide is produced. The hydrogen peroxide can be detected and quantified by horseradish peroxidase (HRP), which reduces the produced H₂O₂ and thereby oxidizes its chromogenic substrate *o*-dianisidine via consecutive one-electron transfers. The oxidized diimine form of *o*-dianisidine can then be measured photospectrometrically [9].

The methodology employs a coupled photospectrometric assay using GOD and HRP with *o*-dianisidine as reporter substrate. The assay was prepared in MTP-format. A reaction solution containing 12.5 U/ml GOD, 2.5 U/ml HRP and 0.125 mg/ml *o*-

dianisidine dihydrochloride in 50 mM sodium acetate pH 5.1 was prepared. Sample solutions from culture supernatants were typically diluted in 9 volumes of water. The reaction was started, by adding 50 µl reaction solution to 25 µl of sample and was incubated at 37 °C and 200 rpm for 30 min in a shaking incubator. 50 µl 6 M sulfuric acid was added to stop the reaction and achieve maximum color development (full oxidation of any *o*-dianisidine charge transfer complexes) (Figure 3.1). The developed pink color was measured at 540 nm in a MTP-reader. A calibration curve of a standard D-glucose solutions (0 to 100 µg/ml), that was always part of the experiments, was used to quantify the sample measurements.

22 3.6.3 In vitro O-methyl transferase (O-MT) assay

O-methyl transferase (*O*-MT) assays were conducted in a total volume of (50 to 100) μl. The standard assay buffer was 100 mM Tris/HCl, 2.5 μM GSH pH 7.5. 1 mM MgCl₂, which was otherwise omitted, was added for reactions using cation dependent *O*-MTs (e.g. PFOMT). Reactions contained 0.5 mM alkyl donor (e.g. (*S*,*S*)-27 SAM) and 0.4 mM flavonoid or cinnamic acid substrate. Enzymatic reactions were started by addition of enzyme (usually 0.2 mg/ml) and incubated at 30 °C.

Reactions were stopped by addition of 500 μl ethyl acetate containing 2 % formic acid and vortexed for 15 s to extract the hydrophobic phenylpropanoids and flavonoids. After centrifugation (10 000 × g, 4 °C, 10 min) the organic phase was transferred into a new tube. The reaction was extraced once more with 500 μl ethyl acetate, oncentrator (Concentrator 5301; eppendorf, Hamburg, Germany). The residue was dissolved in methanol and centrifuged at 10 000 × g for 10 min to remove unsoluble matter. The supernatant was transferred into a HPLC vial and analyzed by HPLC (3.6.8).

BWEIGEL: nochmal auseinanderklamüsern wegen den konzentrationen und eingesetzten enzymmengen...

OMe diamine
$$-e^-,H^+$$
OMe NH₂

$$H_2N$$
OMe NH₂

$$H_1/2$$

$$H_2N$$
OMe NH₂

$$HN$$
OMe NH₂

$$HN$$
OMe NH₂

$$HN$$
OMe NH₃

$$-e^-,H^+$$
OMe NH
$$-e^-,H^+$$
OMe NH
$$HN$$
OMe NH

Figure 3.1.: Oxidation of the reporter substrate o-dianisidine. Consecutive one-electron transfers lead to the fully oxidized diimine form of o-dianisidine. The first electron transfer is believed to produce a charge transfer complex intermediate. [23, 9]

When detection of hydrophobic (e.g. flavonoids) and hydrophilic compounds (e.g. 2 SAM, SAH) was performed simultaneously reactions were stopped by addition of 30.3 volumes 10% (w/v) TCA in 50% acetonitrile. The mixture was vortexed for 30.3 complete mixing and incubated on ice for at least 30 min. After centrifugation 30.3 (10 30.3) (10 30.3) 30.3) 30.30 min) the supernatant was transferred into HPLC-sample vials 30.30 and analyzed (see 3.6.8).

7 Measurement of activity/pH profiles

⁸ Assays to measure activity over larger pH ranges were set up in 50 mM L-malic acid/MES/Tris (MMT)- (pH 4 to 9) or succinate/sodium phosphate/glycine (SSG)¹⁰ buffer (pH 4 to 10) to keep the concentrations of buffer salts constant for each pH [37].

The protein of interest was first extensively dialyzed against the reaction buffer (e.g. MMT, SSG) at pH 7 with added EDTA (5 mM) and then against the same buffer without EDTA. Standard reaction conditions were 50 mM buffer, 0.4 mM salkyl acceptor (e.g. caffeic acid), 0.5 mM SAM, 2.5 μM GSH and 0.2 mg/ml enyzme. MgCl₂ was either omitted or added at 10 mM to assess influences of divalent cations. Assays were stopped as described in 3.6.3 and analyzed accordingly.

18 3.6.4 Photospectrometric assay for the methylation of catecholic moieties

²⁰ Catecholic moieties can form stable complexes in the presence of heavy metals such as copper or iron [48, 34]. Hence, caffeic acid can complex ferric (Fe³⁺) ions ²² and form a colored complex with λ_{max} = 595 nm [12]. Since the complex formation ²³ is specific for caffeic acid and methylated derivatives (i.e. ferulic and iso-ferulic ²⁴ acid) cannot complex Fe³⁺, this can be used as a measure for methylation reactions. ²⁵ *O*-MT assays were prepared as before (3.6.3). However, the reactions were stopped ²⁶ by addition of 0.1 volumes 1 M Tris/HCl pH 8, immediately followed by 0.5 volumes ²⁷ catechol reagent (2 mM FeCl₃ in 10 mM HCl). The complex formation reaction was ²⁸ allowed to equilibrate for 5 min at RT and the absorbance at 595 nm was measured.

29 3.6.5 Hydrophobic interaction chromatography (HIC)

3.6.5 Hydrophobic interaction chromatography (HIC)	BWEIGEL: complete
3.6.6 Analytical gel filtration	
	BWEIGEL: complete

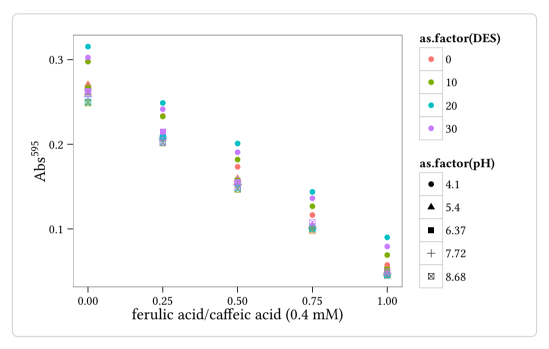


Figure 3.2.: Calibration curves of different relative compositions of ferulic acid to caffeic acid, that were taken as described in 3.6.4. The total concentration was always 0.4 mM. At lower pH values around 4, the method seems to overestimate the concentration of caffeic acid. However, the slope of the curves stays the same.

13.6.7 Binding experiments using Isothermal Titration Calorimetry (ITC)

Isothermal Titration Calorimetry (ITC) can be used to directly characterize the thermodynamics of an observed process, be this a binding interaction or an enzymatic reaction [17].

⁶ ITC measurements to describe the interaction between PFOMT and its substrates/- effector where performed using a MicroCal iTC200 device (Malvern, Worcestershire, ⁸ UK). PFOMT protein was extensively dialyzed against 50 mM MMT-buffer pH 7 prior to ITC experiments. The solution was susequently centrifuged (14 000 × g, ¹⁰ 4 °C, 10 min), to remove insoluble matter and aggregates. The dialysate was stored at 4 °C and used to prepare substrate and effector solutions. Generally 50 μM protein was provided in the ITC cell and the effectors/substrates to be titrated were loaded into the syringe. The substance concentration in the syringe was ten times higher than the protein solution. Experiments were carried out at 20 °C unless otherwise stated. The stirring speed was set to 500 rpm. The injection volume was set to (2 to 4) μl, amounting to a total of 10 to 19 injections.

17 3.6.8 High-performance liquid chromatography (HPLC) analytics

¹⁹ Due to their aromaticity, methanolic extracts of flavonoids exhibit two major

absorption peaks in the UV/VIS region of the light spectrum in the range of (240 to 400) nm [30]. However, even the more simple phenyl propanoids (e.g. cinnamic acids) show absorption of light in the UV/VIS-region.

Methanolic extracts of flavonoids and phenyl propanoids were analyzed by HPLC using a photo diode array (PDA)-detector, which was set to record in the range of (200 to 400) nm. HPLC runs were performed on a reverse-phase C-18 end-capped column (YMC-Pack ODS-A; YMC Europe, Dinslaken, Germany) with a pore size of 120 Å. The mobile phase was aqueous acetonitrile supplemented with 0.2 % formic acid. The flow was kept constant at 0.8 ml/min. 10 μl *O*-MT enzyme assay extract (3.6.3) were injected and analyzed using an acetonitrile gradient starting with 5 % acetonitrile (4 min). The acetonitrile content was increased to 100 % in 21 min and

33 3.6.9 liquid chromatography coupled mass-spectrometry (LC/MS) measurements

31 was kept at 100 % for 5 min. Peaks were integrated from the 280 nm trace using the

32 software provided by the manufacturer of the device.

- 4 Evaluation of PFOMT towards
 the acceptance of long-chain SAM
 analogues
- 44.1 Introduction
- **54.2 Substrate binding studies using ITC**
- 4.3 Determination of the structure of apo-PFOMT
- 74.3.1 PFOMT activity in deep eutectic solvents (DES) / Solubility-
- **enhancing effects of DES**
- y vielleicht eigenes kapitel DES?
- **4.4 Study of variants for long-chain alkylations**
- 11 4.4.1 PFOMT-Paper (DIM)
- 12 4.4.2 **Dockings???**
- 13 4.5 Colclusion/Discussion

5 Enzymatic methylation of Noncatechols

35.1 Introduction

⁴ Non-catechols in nature (biosynthesis, mode of action?), chemical methylation???

- 5.2 **SOMT-2**
- 5.2.1 In vivo methylation studies using N. benthamiana
- 75.2.2 In vivo studies in *E. coli*
- **5.2.3** In vitro studies using recombinantly produced SOMT-2
- **5.3 PFOMT**
- **5.3.1** Acidity and Nucleophilicity of phenolic hydroxyl-groups
- **11 5.3.2 pH-Profiles of PFOMT-catalysis**
- 12 5.3.3 Influence of Mg²⁺ on PFOMT activity
- 5.4 Consensus or Bioinformatic points-of-view(COMT)???
- 15 5.5 Conclusion/Discussion

- Development of an whole cell methyl transferase screening system
- 46.1 Introduction
- **56.2** Theoretical considerations / design of system
- 6.3 Detectability of S-adenosyl-L-homocysteine (SAH)

8 SAM

- 56.4 Usage of the Isr-promoter for true autoinduction
- **...6.5** Conclusion/Discussion

7 DES in protein crystallography

- ₂ 7.1 Introduction
- 37.2 Solubility enhancement of hydrophopbic sub-
- stances by addition of DES
- **5 7.3 Enzymatic** *O***-methylation in DES**
- **7.4 DES** as precipitants in protein crystallization
- 7.5 Conclusion/Discussion

8 Acknowledgements

² Quisque ullamcorper placerat ipsum. Cras nibh. Morbi vel justo vitae lacus tincidunt ³ ultrices. Lorem ipsum dolor sit amet, consectetuer adipiscing elit. In hac habitasse ⁴ platea dictumst. Integer tempus convallis augue. Etiam facilisis. Nunc elementum ⁵ fermentum wisi. Aenean placerat. Ut imperdiet, enim sed gravida sollicitudin, ⁶ felis odio placerat quam, ac pulvinar elit purus eget enim. Nunc vitae tortor. Proin ⁷ tempus nibh sit amet nisl. Vivamus quis tortor vitae risus porta vehicula.

Fusce mauris. Vestibulum luctus nibh at lectus. Sed bibendum, nulla a faucibus semper, leo velit ultricies tellus, ac venenatis arcu wisi vel nisl. Vestibulum diam. Aliquam pellentesque, augue quis sagittis posuere, turpis lacus congue quam, in hendrerit risus eros eget felis. Maecenas eget erat in sapien mattis portitor. Vestibulum porttitor. Nulla facilisi. Sed a turpis eu lacus commodo facilisis. Morbi fringilla, wisi in dignissim interdum, justo lectus sagittis dui, et vehicula libero dui cursus dui. Mauris tempor ligula sed lacus. Duis cursus enim ut augue. Cras ac magna. Cras nulla. Nulla egestas. Curabitur a leo. Quisque egestas wisi eget nunc. Nam feugiat lacus vel est. Curabitur consectetuer.

III Appendix

A Figures

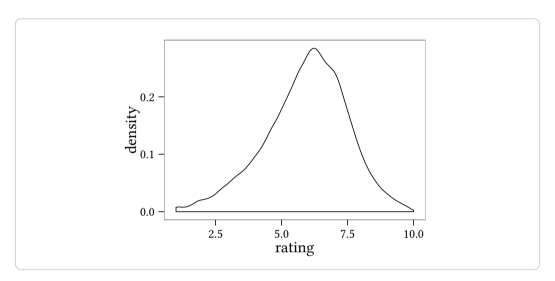


Figure A.1.: Lorem ipsum dolor sit amet, consectetuer adipiscing elit. Aenean commodo ligula eget dolor. Aenean massa. Cum sociis natoque penatibus et magnis dis parturient montes, nascetur ridiculus mus. Donec quam felis, ultricies nec, pellentesque eu, pretium quis, sem.

B Tables

Table B.1.: Overview over the contructs produced for the present thesis. Each step during the production of the construct is given in

construct name	description	entry constructs	destination	workflow steps (primers/cloning sites)
pBEW101	lsrA nromoter			
pBEW103	pBEW102 with BamHI cloning site	pBEW102		
pBEW104	$^{\circ}$ rha $^{ m BAD}$ promoter	pBEW4b	pBEW103	amplification (pRha1.fw/rv), cloning (BgIII,
pBEW106	pICH413038-somt	pET28MC-somt	pICH413038	amplification (somt1/2/3/4), golden gate
pBEW107		pICH51266, pBEW106,	pICH75044	ctoning (bpii) golden gate cloning (BsaI)
		pICH41421		
pBEW1a				
pBEW2a				
pBEW2b				
pBEW3a pBEW3b				
pBEW4a				
pBEW4b pET28-pfomt	pfomt gene in pET-28a(+), endogenous Ndel site removed	pQE30-pfomt	pET-28a(+)	mutagenesis (pfomt1.fw/rv), amplification
pET20-somt	N-terminal pelB-tag fusion for periplasmic expression		pET20-b(+)	(pjomtz.fw/rv), cloning (Ndel, EcoKl)
pET28-somt			pET28-a(+)	
pET32-somt	N-terminal TrX-tag fusion		pET-32a(+)	
pET41-somt	N-terminal GST-tag fusion		pET-41a(+)	
pUC19*	added BglII site	pUC19		mutagenesis ($\rho UCI.fw/rv$)
pUCB1	pUC19 derivative with lsrA promoter	Isr-XX-DAS	pUC19*	cloning (Ndel, BgIII)
pUCB1-sfGFP-DAS+4			4	

C Affidavit

I hereby declare that this document has been written only by the undersigned and
without any assistance from third parties. Furthermore, I confirm that no sources
have been used in the preparation of this document other than those indicated in
the thesis itself.
Date: , Location: , Signature:

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Acronyms

² Å Ångström, 0.1 nm

AC-9 anthracene-9-carboxylic acid 20

⁴ **ATP** adenosine triphosphate 20

BisTris 2-[Bis(2-hydroxyethyl)amino]-2-(hydroxymethyl)propane-1,3-diol

B-PER bacterial protein extraction reagent

CCP4 Collaborative Computational Project No. 4 22, 23

CD circulary dichroism 10

COMT catechol O-methyl transferase 13

- 10 **Coot** Crystallographic Object-Oriented Toolkit 22
- 11 **CV** column volumes
- 12 **DoE** design of experiments 18
- ¹³ **DTT** dithiothreitol; (2*S*,3*S*)-1,4-bis(sulfanyl)butane-2,3-diol
- 14 **EDTA** ethylenediaminetetraacetic acid 15, 17, 18, 26
- 15 **FFD** fractional factorial design vii, 18, 19
- 16 **FPLC** fast protein liquid chromatography 17
- 17 **FT** Fourier transformation 22
- 18 **GdmCl** guanidinium hydrochloride
- 19 **GFP** green fluorescent protein 23
- 20 GOD glucose oxidase 24, 47
- $_{\text{21}}$ $\boldsymbol{GSH}~$ glutathione, $\gamma\text{-L-glutamyl-L-cysteinylglycine}$ 18, 24, 26
- 22 **GSSG** glutathione disulfide 18
- ²³ **HEPES** 2-[4-(2-hydroxyethyl)piperazin-1-yl]ethanesulfonic acid
- ²⁴ **HPLC** high-performance liquid chromatography 13, 20, 24, 28
- 25 **HRP** horseradish peroxidase 24

Acronyms Acronyms

- **IB** inclusion body 17, 19
- **IEX** ion exchange chromatography 20
- **IMAC** immobilized metal affinity chromatography
- 4 IPB Leibniz-Institute of Plant Biochemistry
- FIPTG isopropyl-D-thiogalactopyranosid 14, 16, 17
- 6 ITC Isothermal Titration Calorimetry 28, 47
- **LB** lysogeny broth 12, 13, 16
- 8 MES 2-(N-morpholino)ethanesulfonic acid
- MLU Martin-Luther-Universität
- 10 MMT L-malic acid/MES/Tris 6, 28
- □ MR molecular replacement
- 12 MTP micro-titer plate 20-24, 47
- 13 MW molecular weight 15
- 14 MWCO molecular weight cut-off
- 15 NADES natural deep eutectic solvent vii, 7, 21, 45
- 16 NPS nitrogen, phosphate, sulfate buffer
- 17 NTA nitrilo triacetic acid 17
- ¹⁸ **O-MT** O-methyl transferase 24, 26, 28
- 19 PAGE polyacrylamide gel electrophoresis 14, 15, 19, 20
- 20 PBS phosphate buffered saline 14, 19, 23
- ²¹ **PCH** propane-1,2-diol/choline chloride,NADES-mixture 21
- 22 PCR polymerase chain reaction 10, 11
- ²³ **PDA** photo diode array 28
- 24 **PDB** Protein Data Base 22, 23
- ²⁵ **PFOMT** phenylpropanoid and flavonoid O-methyl transferase 13, 16, 21–24, 28, 47
- ²⁶ **PHENIX** Phyton-based Hierarchial Environment for Integrated Xtallography 23
- 27 **PMSF** phenylmethylsulfonylfluoride
- 28 **RT** room temperature
- 29 **SAE** S-adenosyl-L-ethionine, (2S)-2-amino-4-[[(2S,3S,4R,5R)-5-(6-aminopurin-9-
- yl)-3,4-dihydroxyoxolan-2-yl]methyl-ethylsulfonio]butanoat 20, 21
- 31 **SAH** S-adenosyl-L-homocysteine 18, 26
- 32 SAM S-adenosyl-L-methionine 20, 24, 26, 31
- 33 **SAMS** S-adenosylmethionine synthase 20
- 34 **SDS** sodium dodecylsulfate 7, 14, 15, 19, 20
- 35 **SOMT-2** soy O-methyl transferase 13, 16–20

Acronyms Acronyms

SSG succinate/sodium phosphate/glycine 7

² **TB** terrific broth 13 ³ **TCA** trichloro acetic acid 14, 15, 19, 26 ⁴ **Ti-plasmid** tumor inducing plasmid 8, 47 ⁵ **Tris** tris(hydroxymethyl)-aminomethane

 $_6$ U enzyme unit; measure for enzymatic activity (1 U = 1 µmole/min = 1/60 µkat) $_7$ UV/VIS ultra violet/visible (light spectrum) 20, 28

 $_8$ **V** volume

g **ZYP** N-Z-amine, yeast extract, phosphate 16, 47

Glossary

2 GOD Glucose oxidase is an enzyme.... 44

s Isothermal Titration Calorimetry (ITC) Fill in description here 45

⁴ MTP Micro-titer plate. Small format rectangular plastic plate containing wells to allow for storage of multiple small samples or the containment multiple simultaneous reactions. Typical sizes include 24, 96 and 384-wells 45

PFOMT Phenylpropanoid and flavonoid O-methyl transferase from *Mesembryan-themum crystallinum*, which was first described by Ibdah et al. in 2003 [22]

Ti-plasmid Commonly found plasmids in *A. tumefaciens* and *A. rhizogenes* that confer virulence 46

T2 **ZYP-5052** Autoinduction medium developed by Studier [51]. The naming stems from the components N-*Z*-amine, *y*east extract and *p*hosphate. The numbering designates the composition; e.g. 5052 refers to 0.5 % glycerol, 0.05 % glucose and 0.2 % lactose. 46