Week 10: 10. August 2015- 14. August 2015

10. August 2015

- 1) Digest of hps with Ncol and EcoRI for cloning into pET-28
 - Pipetting scheme according to the protocol Restriction Digest
 - Incubate the digest again for 1 hour at 37 °C
 - Verify 10 µl of the digestion on agarose gel

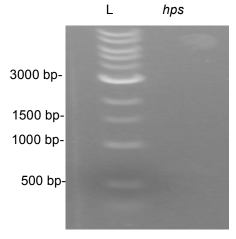


Figure 1: Digest of *hps* **with Ncol/EcoRI.** 10 μl of Digest were checked on 1% (w/v) agarose gel. Expected sizes: *hps*-671 bp. As ladder (L) 1kB Ladder (NEB) was used.

- 2) Digest of hps, phi and medh2 with EcoRI/ PstI for cloning into pSC1B3
 - Pipetting scheme according to the protocol Restriction Digest
 - Incubate the Reaction for 1 hour at 37 °C
 - Verify 10 µl of the digestion on agarose gel

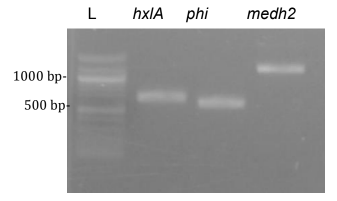


Figure 2: Digest of *hps, phi* and *medh2* with EcoRI/ PstI for cloning into pSC1B3. 10 µI of Digest were checked on 1% (w/v) agarose gel. Expected sizes: *hps*-671 bp, phi-591 bp, *medh2*-1203 bp. As ladder (L) 100bp Ladder (NEB) was used.

- 3) Purification of digested hps, phi and medh2
 - Using Wizard® SV Gel and PCR Clean-Up System (Promega) and following the provided manual
- 4) Ligation of hps, phi and medh2 into pSC1B3 using T4 DNA Ligase
 - Pipetting scheme according to the Protocol Ligation with T4 DNA Ligase (NEB)
 - Use dephosphorylated pSC1B3
 - Control Reaction: pSC1B3 Selfligation
 - Incubate the reaction at 16 °C overnight

11. August 2015

- 1) Digest of hps with Ncol/ EcoRI for cloning into pET-28
 - Pipetting scheme according to the protocol Restriction Digest
 - Incubate the digest again for 1 hour at 37 °C
 - Verify 10 μl of the digestion on agarose gel

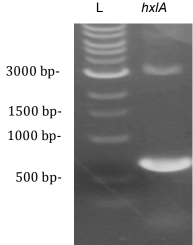


Figure 3: Digest of *hps* **with Ncol/EcoRI.** 10 μl of Digest were checked on 1% (w/v) agarose gel. Expected sizes: *hps*-671 bp. As ladder (L) 1kB Ladder (NEB) was used.

2) Purification of digested hps

- Using Wizard® SV Gel and PCR Clean-Up System (Promega) and following the provided manual
- 3) Ligation of hps and medh2 into pET-28
 - Pipetting scheme according to the Protocol Ligation with T4 DNA Ligase (NEB)
 - Use dephosphorylated pET-28-EcoRI/XhoI (for medh2) and pET-28-NcoI/EcoRI (for hps)
 - Control Reaction: pET-28 Selfligations

- Incubate the reaction at 16 °C overnight
- 4) Transformation of *E. coli* TOP10 cells with pSC1B3+hps, pSC1B3+phi and pSC1B3+medh2
 - Add 5 µl of Ligation reaction to chemically competent *E. coli* TOP10 cells.
 - Following the Protocol for Transformation of chemical competent *E. coli* cells
 - Plate on LB+Clm [25 µg/ml] and incubate overnight at 37 °C

12. August 2015

- 1) Colony-PCR screen for *medh2*, *hps* and *phi* in pSC1B3
 - Constructs: pSC1B3+hps, pSC1B3+phi and pSC1B3+medh2
 - Pipetting scheme and PCR program according to PCR with Taq-DNA Polymerase Protocol
 - Primer: medh2-BioBrick Fwd/Rev, phi-BioBrick Fwd/Rev, hps-BioBrick Fwd/Rev
 - Check 6 Clones
 - Positive Control: add 1 µl of pSC1B3+mmoB
 - Negative Control: add 1 µl MilliQ Water

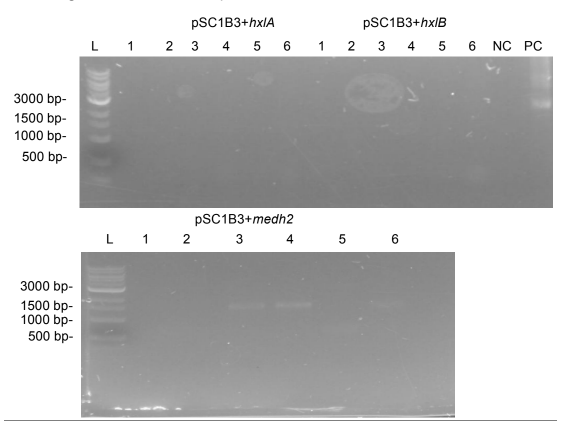


Figure 4: Colony-PCR to identify clones containing pSC1B3+hps, pSC1B3+phi and pSC1B3+medh2 Biobrick constructs.

PCR with Taq-DNA Polymerase to identify pSC1B3+hps, pSC1B3+phi and pSC1B3+medh2. Numbers 1-6 determine the checked clone. As positive control (PC) 1 µl pCR4+medh2 (A) and pSC1B3+mmoB (B) was used. As negative control (NC) 1 µl MilliQ Water was added. 10 µl of PCR were analyzed on 1 % (w/v) agarose gel. Expected sizes: hps-671 bp, phi-591 bp, medh2-1203 bp. As ladder (L) 1 kB Ladder (NEB) was used.

- The colony-PCR failed, nevertheless we inoculated some cultures for plasmid isolation.
- 2) <u>Inoculation of liquid culture for plasmid isolation of pSC1B3+hps, pSC1B3+phi</u> and pSC1B3+medh2
 - Inoculate clone with 5 ml LB+Clm [25 μg/ml] and incubate overnight at 37 °C shaking at 220 rpm.
- 3) <u>Transformation of *E. coli* TOP10 cells with pET-28+*hps*, pET-28+*medh2* and pET-28 Selfligation</u>
 - Add 5 µl of Ligation reaction to chemically competent *E. coli* TOP10 cells.
 - Following the Protocol for Transformation of chemical competent E. coli cells
 - Plate on LB+Clm [25 µg/ml] and incubate overnight at 37 °C

13. August 2015

- Ligation of hps and medh2 into pET-28 failed
- 1) Plasmid Isolation of pSC1B3+hps, pSC1B3+phi and pSC1B3+medh2
 - Use the QIAprep Spin Miniprep Kit (Qiagen) and follow the provided manual.
 - Send the plasmids for sequencing with BioBrick Sequencing Primer FWD/Rev
 - Sequencing revealed 100 % integrity of our BioBrick Constructs