

Experiment Design for reconstruction of pTAL1 in GGA

## DAY1

1. Choose proper promoter, RBS and terminator for the vector.

Promoter and RBS:

(1) Constitutive

BBa\_K081005: [http://parts.igem.org/Part:BBa\\_K081005](http://parts.igem.org/Part:BBa_K081005)

Spring 2014 Distribution

16E

201

(2) Inducible

BBa\_K081005+**BBa\_C0040**+BBa\_B0015+BBa\_K081005

BBa\_C0040: [http://parts.igem.org/wiki/index.php?title=Part:BBa\\_C0040](http://parts.igem.org/wiki/index.php?title=Part:BBa_C0040)

Spring 2014 Distribution

2P

Terminator:

BBa\_B0015: [http://parts.igem.org/wiki/index.php?title=Part:BBa\\_B0015](http://parts.igem.org/wiki/index.php?title=Part:BBa_B0015)

Spring 2014 Distribution

3F

2014

TetR repressible promoter

BBa\_R0040: [http://parts.igem.org/Part:BBa\\_R0040](http://parts.igem.org/Part:BBa_R0040)

Spring 2014 Distribution

6F

201

2. Assemble method: Assembly standard 10

[http://parts.igem.org/Assembly\\_standard\\_10](http://parts.igem.org/Assembly_standard_10)

3. Transform the selected plasmids and plate them onto specific antibiotics-containing plate.

4. Design primers for the wanted part of pTAL.

Forward:

IG14003-pTAL-for:

ATTGAATTCGCGGCCGCTTCTAGATGGATCCCATTTCGTCCGC

Reverse:

IG14002-pTAL-re:

TATCTGCAGCGGCCGCTACTAGTATTTCACTGAGGCAATAGCT

## DAY2

1. Pick up several colonies from the plates and incubate them for 6 hours.
2. Extract plasmids and run a gel or sequence it for further confirmation (day3).  
Enzyme digestion system:

plasmids      10 $\mu$ l

ddwater      7 $\mu$ l

CutSmart buffer    2 $\mu$ l

EcoRI      0.5 $\mu$ l    For B0015, use EcoRII and Esp3I

SpeI      0.5 $\mu$ l    For K081005, use NheI and Esp3I

Incubate in 37°C for 2h.

Expected results: K081005: 0.6k+0.4k; B0015: 0.6k + 1.5k

3. After getting the synthesized oligo for the primers, use them to get PCR products of pTAL.

The PCR reaction system (50 $\mu$ l):

pTAL 1 $\mu$ l (20ng)

dNTP 1 $\mu$ l (10mM)

Primer1 (IG14003-pTAL-for) 2.5 $\mu$ l (10nM)

Primer2 (IG14003-pTAL-for) 2.5 $\mu$ l (10nM)

Phusion Polymerase      0.5 $\mu$ l

Phusion Reaction Buffer (HF & GC) 10 $\mu$ l

(another 1.5 $\mu$ l DMSO in GC buffer)

ddwater 32.5 $\mu$ l

Reaction protocol:

(1) 95°C      4min

- (2) 95°C 30s
- (3) 55°C 30s
- (4) 72°C 90s
- (5) Goto Step 2 for 30 cycles
- (6) 72°C 10min
- (7) 4°C Forever

Reference: <https://www.neb.com/protocols/1/01/01/pcr-protocol-m0530>

4. Run a gel for the PCR products (2k) and if the result is positive, sequence it.

## DAY3

1. Use EcoRI and SpeI to digest the PCR products and EcoRI and XbaI to digest BBa\_B0015.  
Also, use EcoRI and SpeI to digest the PCR products and BBa\_B0015 respectively.

Enzyme Digestion System:

PCR products 10µl/5µl

ddwater 7µl/12µl

CutSmart buffer 2µl

EcoRI 0.5µl

SpeI/XbaI 0.5µl

Incubate in 37°C for 2h.

2. Use DNA clean kit to pure the digestion product.

3. Ligation:

PCR products: backbone = 3:1

Quick ligase buffer 5µl

Quick ligase 0.5µl

ddwater

Incubate at 37°C for 5min.

4. Transformation and Screening (Chl-antibiotics)

5. Use the same method to ligate BBa\_C0040 and BBa\_B0015.

## DAY4

1. Pick up several colonies and culture them in chl-containing culture.

## DAY5

1. Extract the reconstructed plasmids (TAL + terminator).

2. Enzyme digestion and run a gel.

Enzyme Digestion System:

plasmids      5 $\mu$ l

ddwater      12 $\mu$ l

CutSmart buffer    2 $\mu$ l

EcoRI      0.5 $\mu$ l

SpeI      0.5 $\mu$ l

Incubate at 37°C for 2h.

Expected result: 1k: 3k.

3. Sequence it.

## DAY6

If the result is positive, continue to construct the plasmid.

1. Use EcoRI and SpeI to digest the ligated products and EcoRI and XbaI to digest BBa\_Q04400/BBa\_K081005.

2. Run a gel to separate wanted bands from digestion products (TAL + terminator.)

3. Use DNA clean kit to pure the digestion product of Use DNA clean kit to pure the digestion product.

4. Ligation.

PCR products: backbone = 3:1

Quick ligase buffer 5 $\mu$ l

Quick ligase 0.5 $\mu$ l

ddwater

Incubate at 37°C for 5min.

5. Transformation and Screening (for K081005, chl)

6. Use the same method to ligate BBa\_C0040+BBa\_B0015 and BBa\_K081005).

## DAY7

1. Extract the plasmids.

2. Enzyme digestion and run a gel.

Enzyme Digestion System:

plasmids 5 $\mu$ l

ddwater 12 $\mu$ l

CutSmart buffer 2 $\mu$ l

EcoRI 0.5 $\mu$ l

SpeI 0.5 $\mu$ l

Incubate at 37°C for 2h.

Expected results: 2k: 3k.

3. Sequence it.

