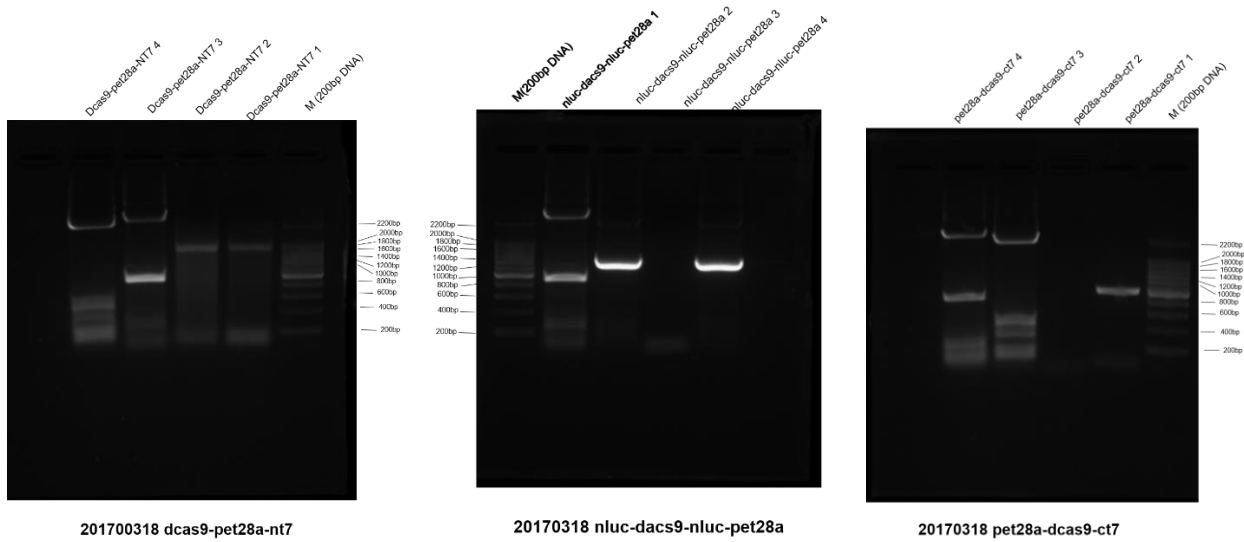


Mar.18<sup>th</sup>

**Q5 PCR** amplified NFluc and CFluc from pBEST-Fluc; NT7 and CT7 from BL21-DE3; dCas9 from px335. We also got pet28a plasmid



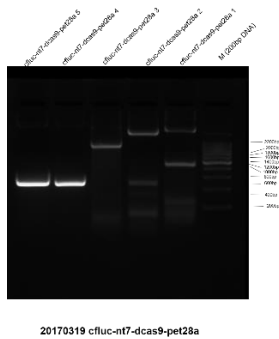
**Assembled, transformed Gibson** into competent cells and **incubated** around 37 celcius

- had good result

Mar.19<sup>th</sup>

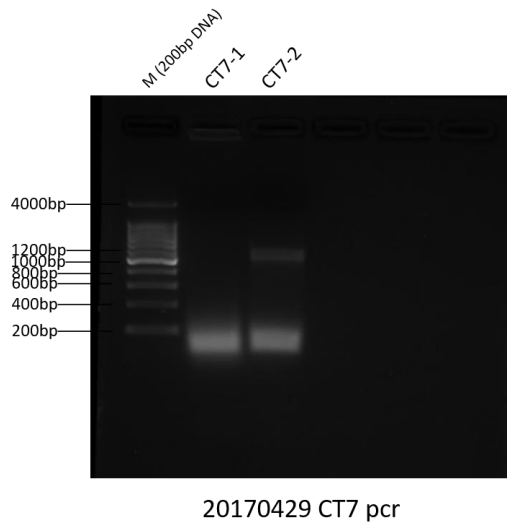
**Enzyme digestion, transformed plasmid** into competent cells and **incubated** around 37 celcius

- NFluc-dCas9
- CFluc-dCas9
- NT7-dCas9



**Apr.29<sup>th</sup>**

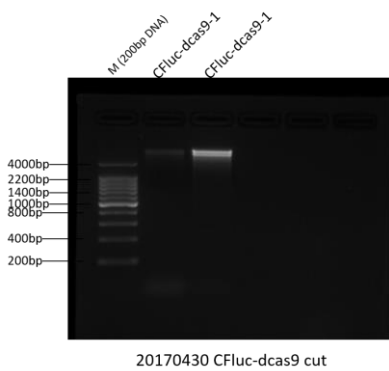
**PCR amplified CT7**



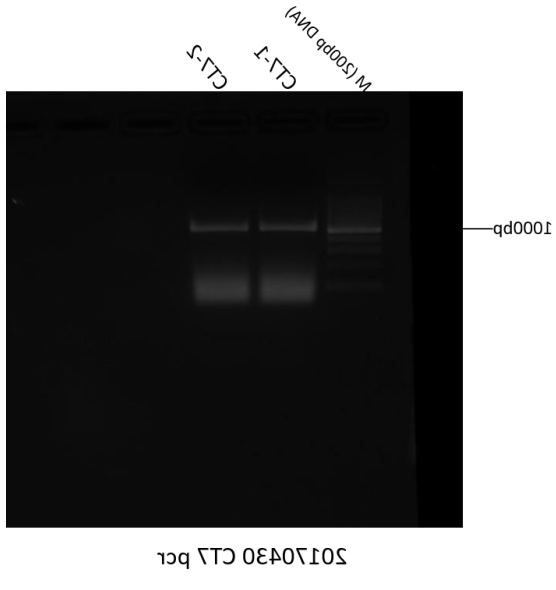
**Apr.30<sup>th</sup>**

**Enzyme digestion**

- CFluc-dCas9
- CT7-dCas9



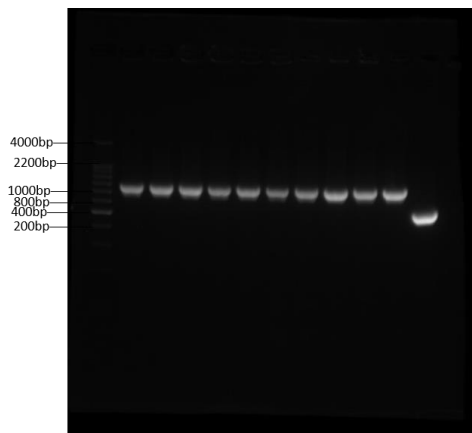
**Ligation of CT7 onto plasmid backbone , transformed plasmid into competent cells and incubated around 37 celcius**



## May.1<sup>st</sup>

**Colony PCR of CT7** proved that we had correct plasmid

- Plasmids
  - pet vr
  - dCas9 vr

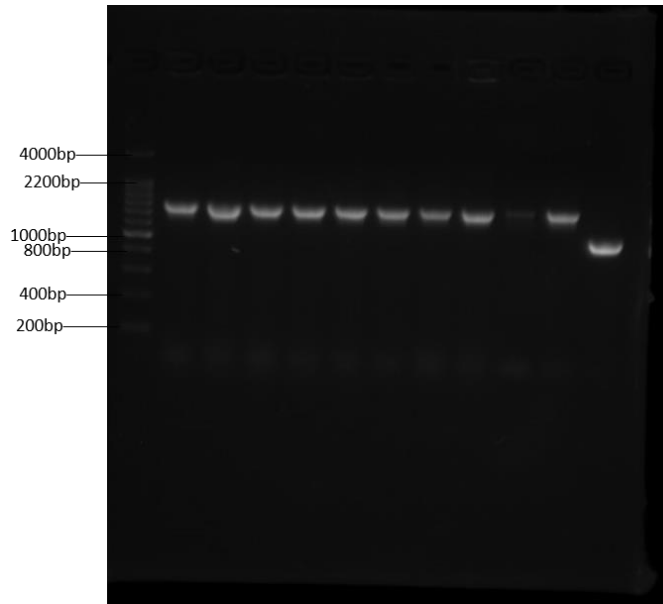


20170501 ct7-dcas9 colony pcr

## May.2<sup>nd</sup>

**Colony PCR of CT7** proved that we had correct plasmid

- Plasmids
  - pet vr
  - dCas9 vr



20170502 colony pcr

---

**Jun.26<sup>th</sup>**

**Construction of 7 sgRNA**

**Diluted primer (F1&FN) into 10 $\mu$ l, 3 $\mu$ l primer was mixed well with DNA to make template**

**First time PCR (20  $\mu$ l)**

Volume:  $\mu$ l

Template	2
DNTPs	1.6
High CG enhancer	1.6
5x Q5 buffer	4
Q5 DNA polymerase	0.2
H2O	10.6

**Reaction time**

Temperature (Celsius)	Time (second)
98	30
98	10
55	1
(Ta) 60	30
72	30
72	2 min

### Second time PCR (50 $\mu$ l) using rTaq system

Volume:  $\mu$ l

#### rTaq

Template (first time PCR product)	10
DNTPs	4
10 x PCR buffer	5
rTaq DNA polymerase	0.5
F1	2
FN (F6)	2
H2O	26.5

#### Reaction time

Temperature (Celsius)	Time (second)
94	5 min
94	30
60	30
72	30
Repeat 2-4	30 cycles
72	2 min
12	$\infty$

Volume:  $\mu$ l

#### Q5 (50 $\mu$ l)

Template	10
dNTPs	4
High CG enhancer	4
5x Q5 buffer	10
Q5 DNA polymerase	0.5
F1	2
FN (F6)	2

H2O	17.5
-----	------

### Reaction time

Temperature (Celsius)	Time (second)
98	30
98	10
60	30
72	15
Repeat 2-4	30 cycles
72	2 min
72	∞

**Jun.28<sup>th</sup>**

### Colony PCR

#### Mix together (800 µl)

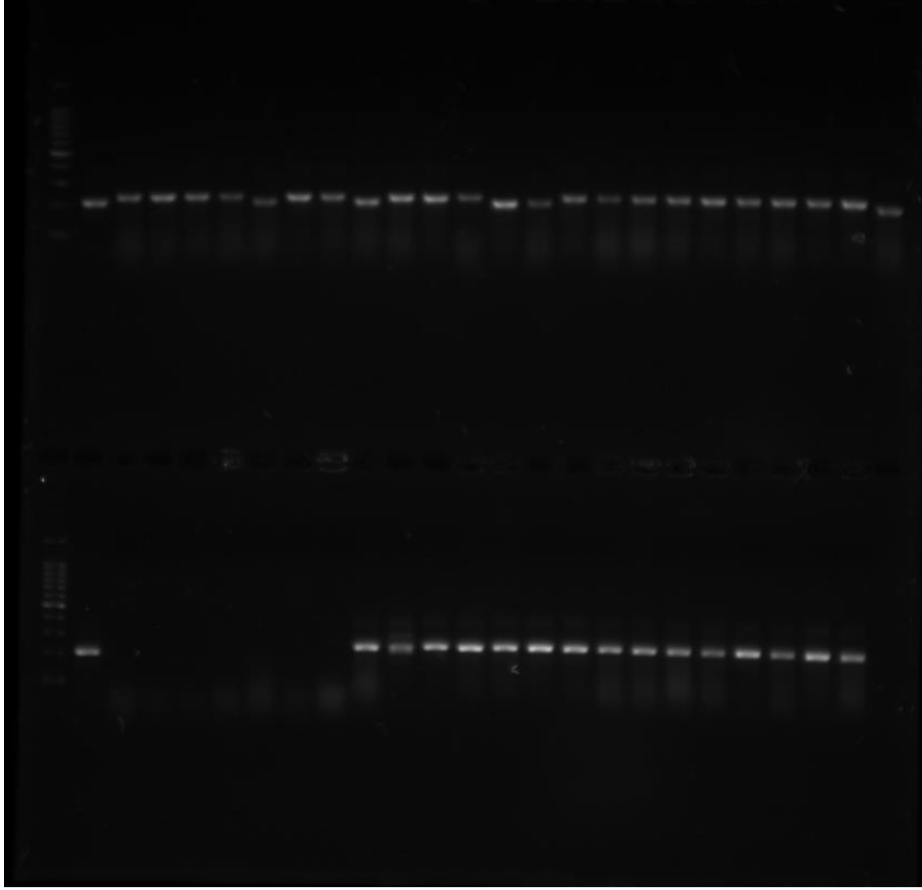
pcr buffer	80
dNTPs	64
VF	32
VR	32
rTaq	8
ddH2O	584

- Make up mix into 1.5 ml PE tube and dispensed into eight pipes
- picked bacteria inside fume hood and put them into those pipes

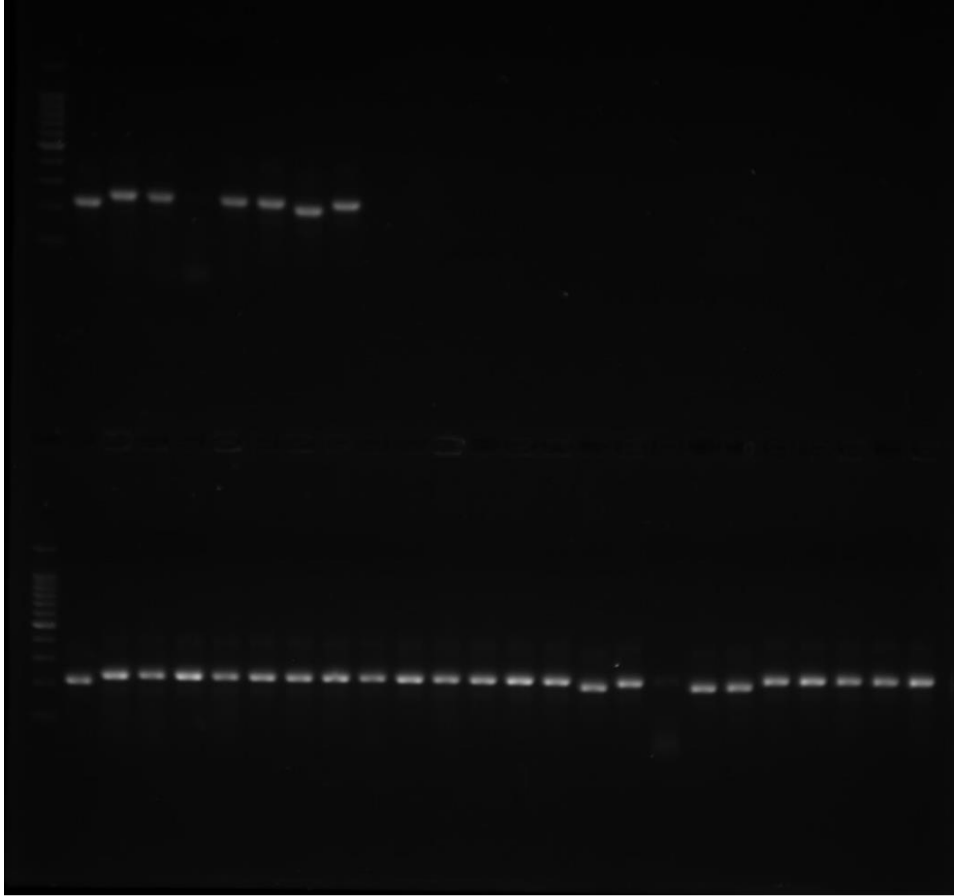
### PCR

#### Reaction time

Temperature (Celsius)	Time (second)
94	5 min
94	30
55	30
72	30
Repeat 2-4	30 cycles
72	5 min
12	∞



2017-6-28 (1,2,3)



2017-6-28 (4,5)

## Protein experiment of CT7 and NT7(without ultrasonic crashing)

Jul.3<sup>rd</sup>

**Transformed plasmid** into BL21 competent cells and **incubated** around 37 celcius

Jul.4<sup>th</sup>

**Inoculation and pre-culture**

Jul.5<sup>th</sup>

**Seed-culture** at 25 Celsius using 5ml tube



- Started with od 0.01 to od 0.5

### **Protein induction I**

- Added different concentration of IPTG

### **Jul.6<sup>th</sup>**

### **Protein induction II**

- After 19 h, collected the bacteria (20ul bacteria plus 5ul loading buffer, 15 minutes under 98 Celsius, lowered the temperature into room temperature and stored inside -20 Celsius refrigerator)

### **Jul.7<sup>th</sup>**

**Made protein gel and gel electrophoresis** to detect if we have the protein or not

- CT7
- NT7

### **Jul.9<sup>th</sup>**

**Made protein gel and gel electrophoresis** to get better gel using same bacteria sample

- CT7
- NT7

### **7.14 分沉淀上清**

### **Jul.10<sup>th</sup>**

### **Enzyme digestion and ligation**

- pSB1C3-va23
- pSB1C3-vb126

## concentration

Plasmid name	Concentration (ng/μl)
pSB1C3	3.4
pSB1C3-va23	12.3
pSB1C3-vb126	8.5

**Transformed plasmid** into BL21 competent cells and **incubated** around 37 celcius (separate precipitation and supernatant)

- CFluc-dCas9-pSB1C3
- NFluc-dCas9-pSB1C3

## Inoculation and pre-culture

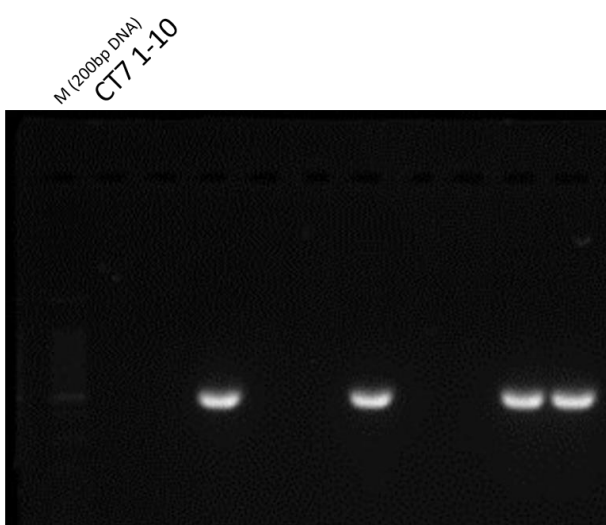
- CFluc-dCas9-pSB1C3
- NFluc-dCas9-pSB1C3

## Ligation

**July 11<sup>th</sup>**

**Restriction Digestion** using BcuI and XbaI

- -PSB1C3



### **Small DNA Fragment Purification** nanodrop

Sample	Concentration (ng/ $\mu$ )
PSB1C3	3.9

- Mistake, instead of transforming and growing more pSB1C3, all of the plasmids were digested. Anyhow, we had more !

### **Transformation** of VA83-PSB1C3, VB126-PSB1C3, PSB1C3

- The plasmid was in an empty tube with a little fluid at the bottom.

### **Overnight cultures**

- NT7-dCas9-PSB1C3 CT7-dCas9-PSB1C3
- Miniprep the other day

### **Glycerol stocks**

- NT7-dCas9-PSB1C3
- CT7-dCas9-PSB1C3

### **Overnight culture**

- CFluc-dCas9-pSB1C3
- NFluc-dCas9-pSB1C3

### **July 12<sup>th</sup>**

### **Made protein gel and gel electrophoresis**

- CT7 1
- CT7 2
- CT73
- CT74
- NT7 1

- NT7 2
- NT7 3
- NT74

## **NFluc, CFluc**

**Seed-culture** at 25 Celsius using 5ml tube

- Started with od 0.01 to od 0.5

### **Protein induction I**

- Added different concentration of IPTG

## **July 13th**

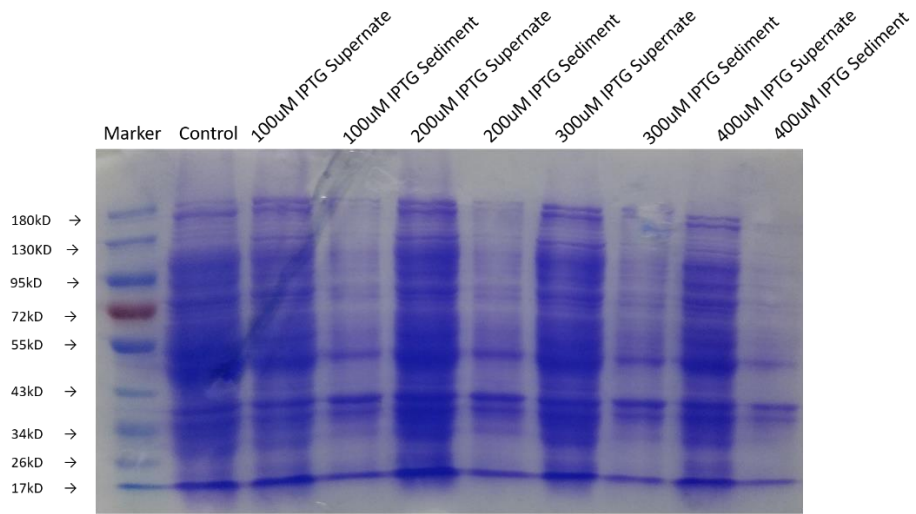
### **Protein induction II**

- After 19 h, collected the bacteria (20ul bacteria plus 5ul loading buffer, 15 minutes under 98 Celsius, lowered the temperature into room temperature and stored inside -20 Celsius refrigerator)

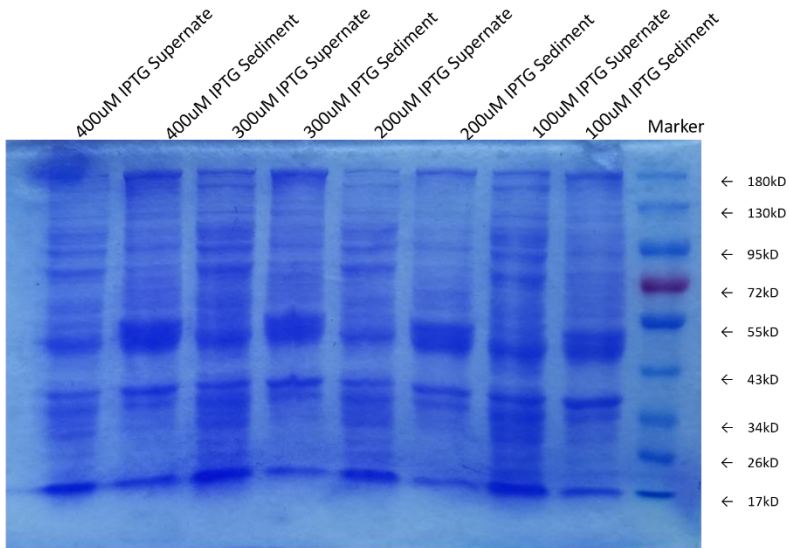
## **July 14<sup>th</sup>**

### **Made protein gel and gel electrophoresis**

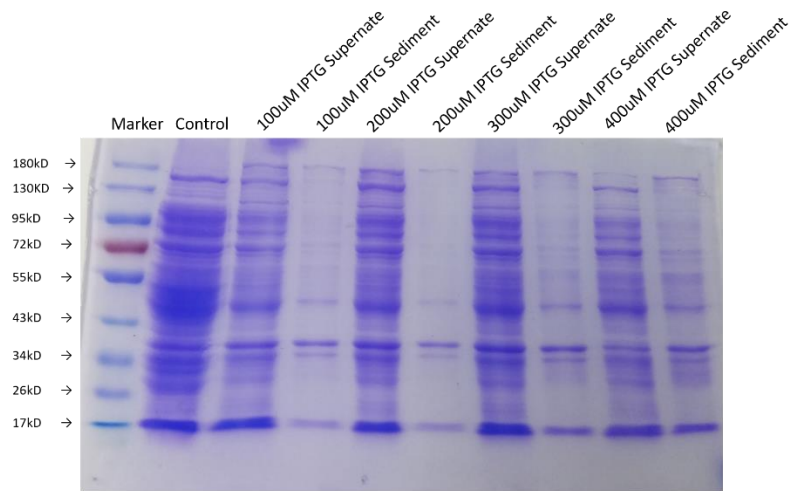
- Order: 0, marker, 0.1nM, 0.2nM, 0.3nM, 0.4nM
- NT7
- CT7
- NFluc
- CFluc



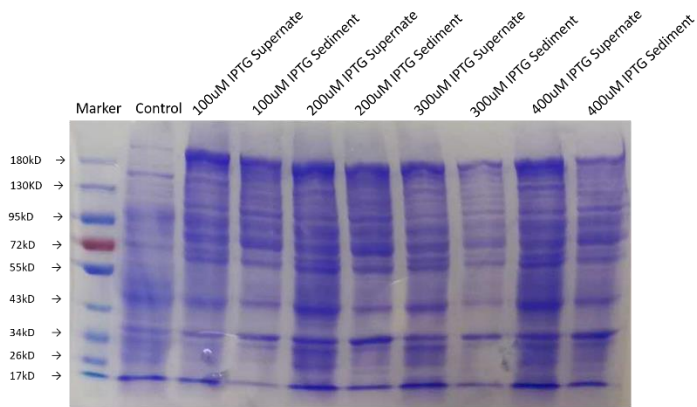
20170714 cflu-dcas9



20170714 CT7-dCas9



20170714 Nfluc-dCas9



20170714 NT7-dCas9

## July 15<sup>th</sup>

### Accidents

- PCR instrument power failure
- Rerun VB126-PSB1C3,VA23-PSB1C3,NT7-PSB1C3,CT7-PSB1C3 colony PCR

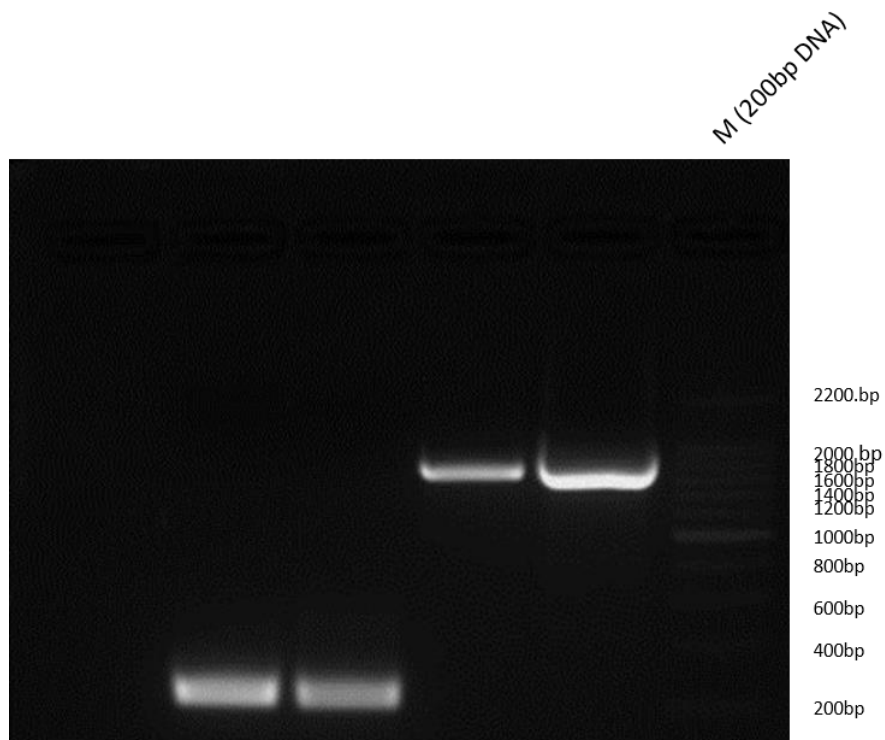
### Gel electrophoresis

- Mistakes: use inappropriate primers

## Possible successfully constructed plasmids

- VA23 3
- VA23 4
- VA23 5
- VA23 6
- VA23 7
- VB126 2
- VB126 7
- VB126 8
- CT7 7
- NT7 8

Re PCR plasmids or digest colony PCR products, gel electrophoresis to testify the other day.



**20170715 VB126 VA23 NT7**

**Prepare selection plates** with Chloramphenicol (Chl)

## July 16<sup>th</sup>

### Made protein gel and gel electrophoresis

#### Protein purification using kit from takara

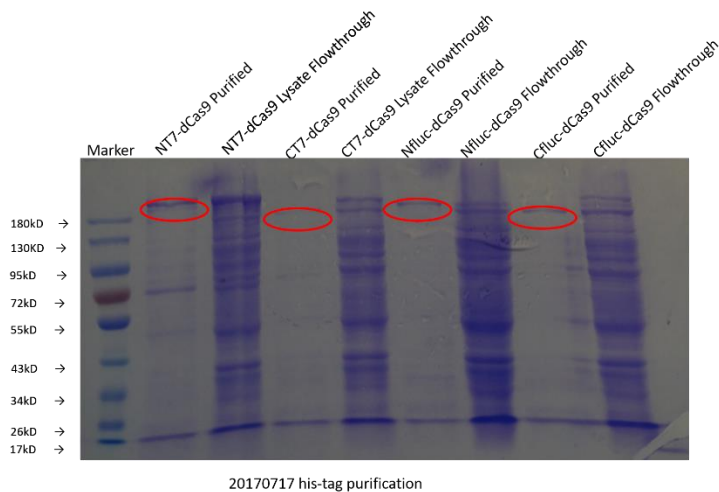
- Keep the supernatant and waste, distributed into 20  $\mu$ l plus 5  $\mu$ l loading buffer, 15 minutes under 98 Celsius, lowered the temperature into room temperature and stored inside -20 Celsius refrigerator

## July 17<sup>th</sup>

### Made protein gel and gel electrophoresis using July.16th product

#### Seed culture (concentration of IPTG 100Nm)

- CT7: 0, 0.05, 0.3
- NT7/NFluc/CFIc: 0,0.05,0.2



## July 20<sup>th</sup>

### Made protein gel and gel electrophoresis using July.16th product

- NFluc 0
- NFluc 1
- NFluc 2



- CFluc 0
- CFluc 1
- CFluc 2
- NT7 0
- NT7 1
- NT7 2

### RNA transcription DNase I

- 33a
- 93a
- 70b
- 147b

### July 21<sup>st</sup>

### Seed culture (concentration of IPTG 100Nm)

- CT7/NT7/NFluc/CFlc: 0,0.03nM,0.06nM,0.09nM,0.2nM(CT7:0.3nM)

#### **IPTG concentration**

nM	μl
0.1	5
0.03	1.5
0.06	3
0.09	4.5
0.2	10
0.3	15

#### **OD**

NT7	1 <sup>st</sup> time	2 <sup>nd</sup> time	3 <sup>rd</sup> time
0	0.12	0.11	0.07
1	0.08	0.08	0.07
2	0.07	0.08	0.07
3	0.10	0.09	0.07
4	0.09	0.11	0.07
CT7			
0	0.09	0.03	0.04
1	0.11	0.09	0.05
2	0.07	0.06	0.09
3	0.09	0.09	0.08

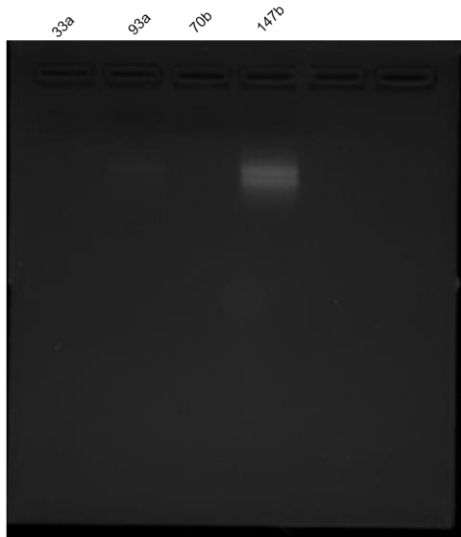
CFluc			
0	0.07	0.07	0.15
1	0.13	0.08	0.14
2	0.09	0.07	0.13
3	0.07	0.15	0.16
NFluc			
0	0.11	0.13	0.20
1	0.06	0.17	0.19
2	0.14	0.09	0.15
3	0.11	0.07	0.19
4	0.08	0.08	0.21

## RNA transcription and gel electrophoresis

- 33a
- 93a
- 70b
- 147b



20170721 第一次RNA电泳 从右到左 33A 93A 70B 147B



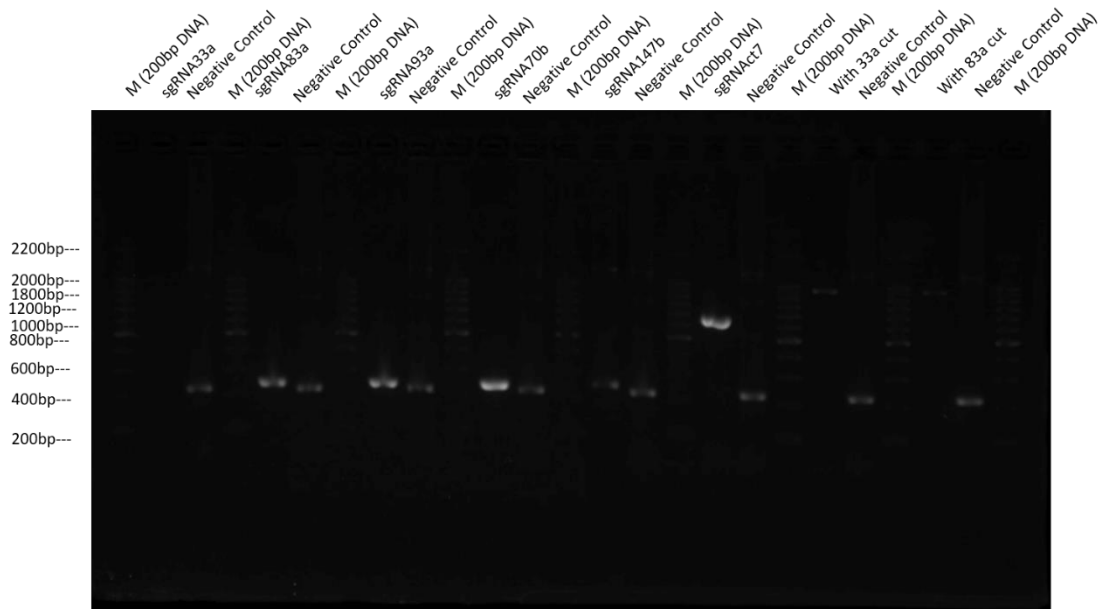
20170721 第二次RNA电泳 33a 93a 70b 147b

**July 23<sup>rd</sup>**

**Restriction digest of plasmid** using Pst1 and EcoR1

- 33a
- 83a
- 93a
- 70b
- 147b
- CT7

**Gel electrophoresis of digestion product**



20170723 33a 83a 93a 70b 147b ct7 pcr with 33a 83a cut

**July 24<sup>th</sup>**

**33a**

**Polymerase chain reaction** using rTaq

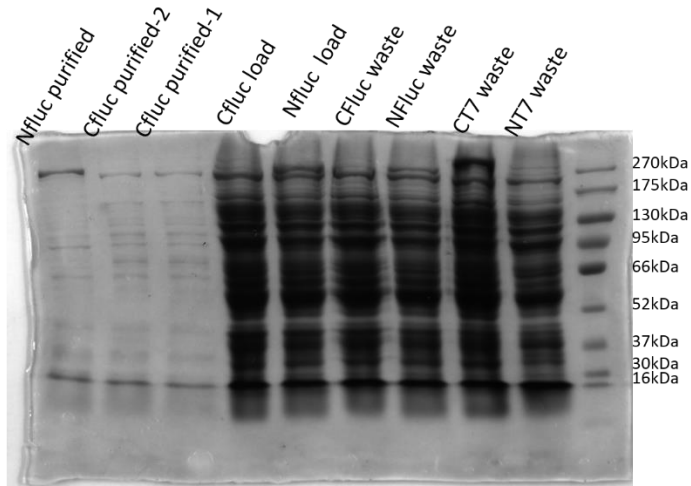
**Restriction digest of plasmid** using Pst1 and EcoR1

**PCR** using Q5

- 23a
- 126b

**Protein ultrasonic crushing**

- NT7
- CT7
- CFluc
- NFluc



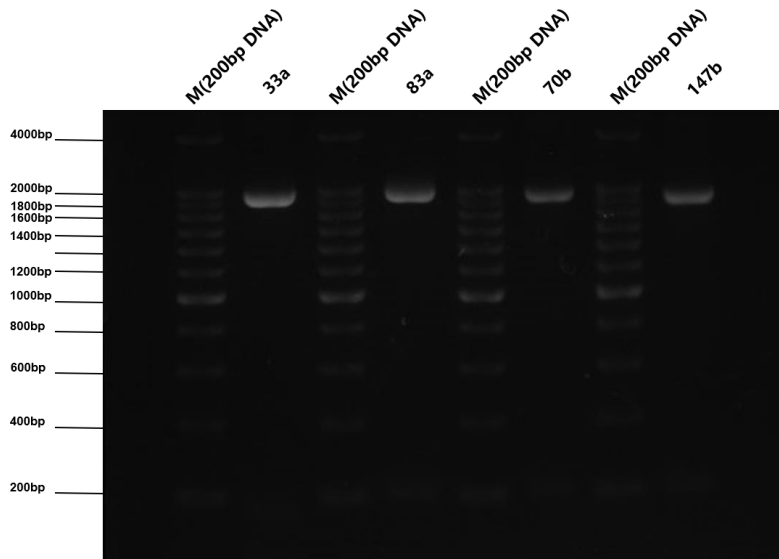
20170724 protein-2 M NT7w CT7w Nfw CFw NFI CFI CFp CFp-2 NFp

## July 25<sup>th</sup>

### Restriction digest of plasmid using Pst1 and EcoR1

- 33a
- 83a
- 70b
- 147b

### Gel electrophoresis of digestion product



20170725 33a 83a 70b 147b RD test

**July 26<sup>th</sup>**

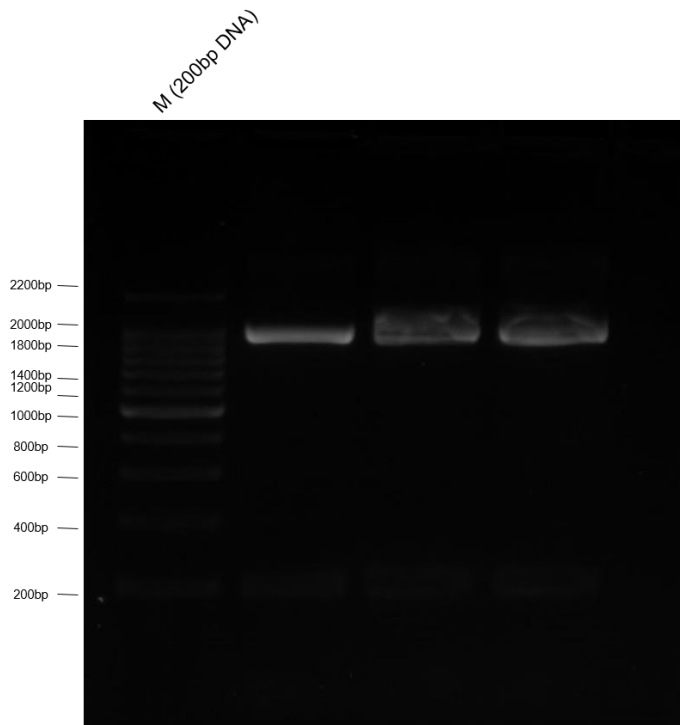
**Preparation for tomorrow's experiment**

**Human practice**

**July 27<sup>th</sup>**

**Gel extraction** dissolve in buffer EB

- 23a
- 126b
- NT7



**20170727 pcr 33a-1c3,6,8,8**

**Restriction digest of plasmid** using BamH1(0.75 $\mu$ ) and Bas1(0.75  $\mu$ )

- CFluc
- CT7
- NT7
- Redo 33a

## July 28<sup>th</sup>

**Colony PCR** no control; extension length for NT7 was too short

- 23a
- 126b
- NT7

**Colony PCR** second time

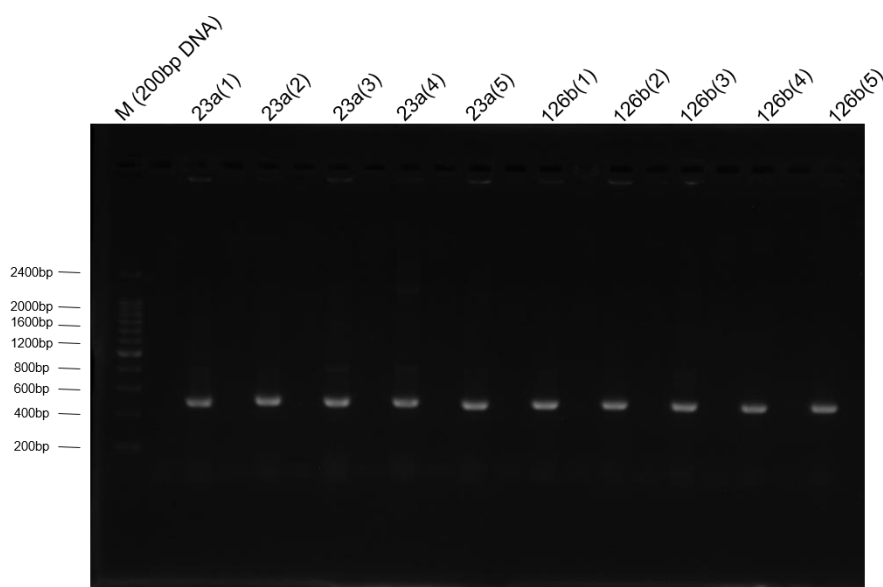
- NT7

## July 30<sup>th</sup>

**Colony PCR** second time, control had some problem

- 23a
- 126b

**Gel electrophoresis NT7** use the wrong enzyme when doing enzyme digestion



2017-7-30 23a12345 126b12345

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## July 31st

## Plasmid extraction and nanodrop

First time

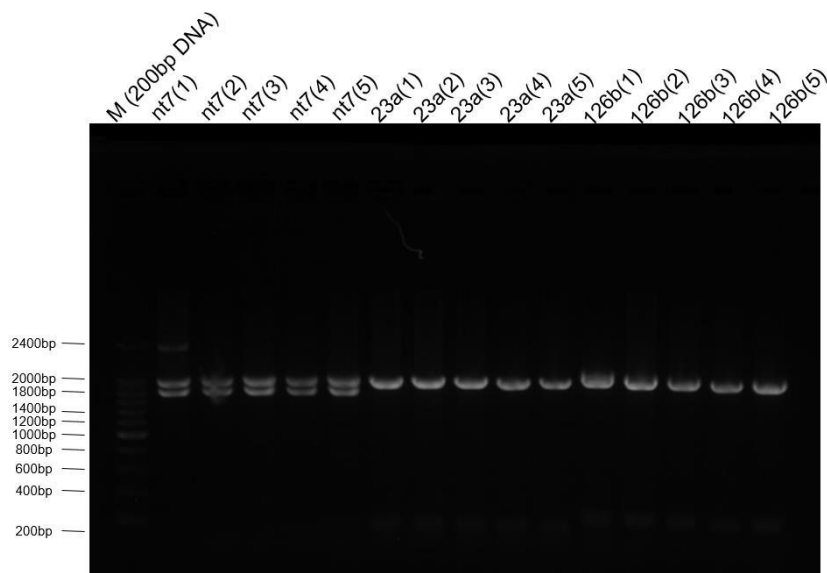
Bacteria sample	1	2	3	4	5
23a	276	255.1	266.3	302.6	273.0
126b	324.4	346.7	379.6	297.4	308.5
NT7	516.4	337.7	368.2	308.5	383.5

Second time

Bacteria sample	1	2	3	4	5
23a	273.3	246.2	267.7	302.1	271.9
126b	321.7	333.3	366.1	286.9	287.2
NT7	540.7	332.3	363.6	298.7	380.0

## Restriction digest of plasmid using Pst1 and EcoR1

### Gel electrophoresis of digestion product



2017-7-31 nt712345,23a12345,126b12345

Aug.1<sup>st</sup>



**Sent the correct plasmid from gel electrophoresis for sequencing**

**Overnight culture** with psb1c3 plasmid backbone

- 33a
- 83a
- 93a
- 70b
- 147b

**Aug.2<sup>nd</sup>**

**Interlaboratory study**

**Transformed dna** into competent cell and **incubated** around 37 celcius

- Positive control
- Negative control
- Device 1
- Device 2
- Device 3
- Device 4
- Device 5
- Device 6

**Aug.3<sup>rd</sup>**

**Tested the sequencing result**, they were all matched,

**Interlaboratory study**

**Transferred the bacteria to the tube**

**Over night culture**

- Positive control
- Negative control
- Device 1
- Device 2
- Device 3
- Device 4
- Device 5
- Device 6

**PCR and gel electrophoresis:** Q5 enzyme was not enough, so we could not see anything in the picture. We threw the gel after.

- 23a
- 33a
- 83a
- 93a
- 70b
- 126b
- 147b

### Aug.4<sup>th</sup>

**Visited the national gene bank and read raw fluorescence** using plate reader

**PCR and gel electrophoresis:** the gel was too light, we threw the gel again and repeat the same experiment after

- 23a
- 33a
- 38a
- 93a
- 70b
- 126b
- 147b

### **Second time PCR**

- 23a
- 33a
- 83a
- 93a
- 70b
- 126b
- 147b

### Aug.5<sup>th</sup>

#### Interlaboratory study

**Redo the transformation and incubation** -none of them grew new bacteria

- Positive control
- Negative control
- Device 1
- Device 2
- Device 3
- Device 4
- Device 5
- Device 6

## RNA transcription, purification, nanodrop and gel electrophoresis

- 23a
- 33a
- 83a
- 93a
- 70b
- 126b
- 147b



**Aug.6<sup>th</sup>**

**Interlaboratory study**

**Transform of PC, NC, D1, D2,D3,D4,D5,D6**

- Increased the time of culture
- Added bacteria medium without antibiotic

**Aug.7<sup>th</sup>**

**RNA transcription**

- 23a
- 33a
- 83a
- 93a
- 70b
- 126b
- 147b

**Aug.8<sup>th</sup>**

**Found out some mistakes of our design**

**Design primer**

**August.9<sup>th</sup>**

**Interlaboratory study**

**Read raw fluorescence** using plate reader (practice)

**Inoculated the bacteria**

**August.10<sup>th</sup>**

**Interlaboratory study**

**Inoculated the bacteria**

Read raw fluorescence until 2 hours

**August.11<sup>th</sup>**

Made protein gel and gel electrophoresis

protein purification, induction and ultrasonic crushing

- NFluc
- CT7

Interlaboratory study

Inoculated the bacteria

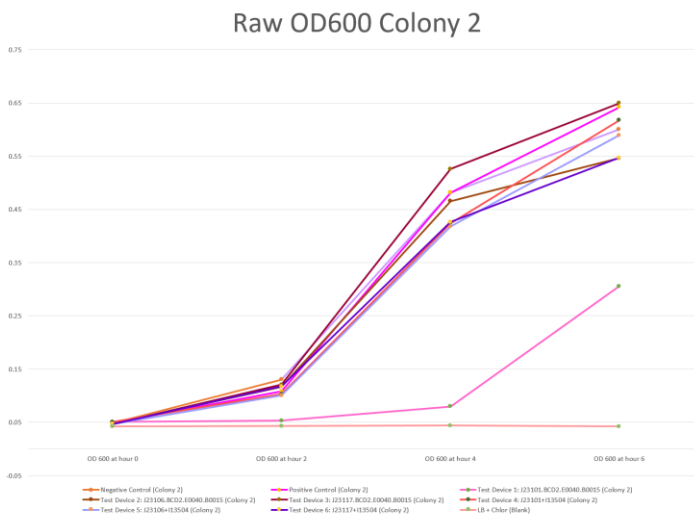
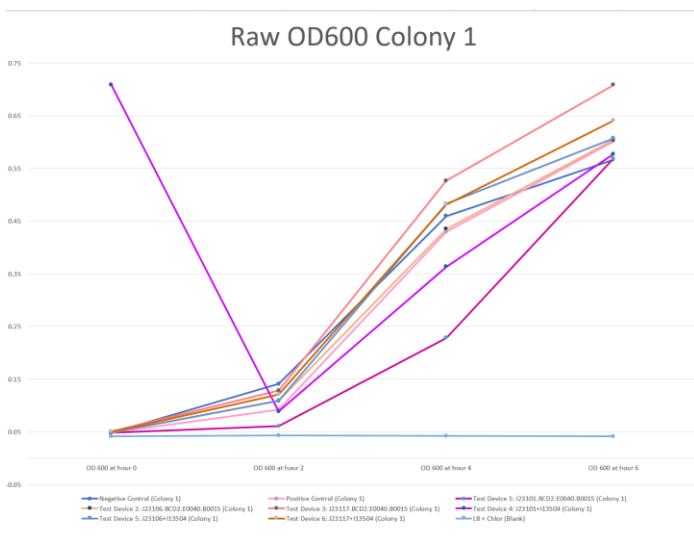
Read raw fluorescence using plate reader

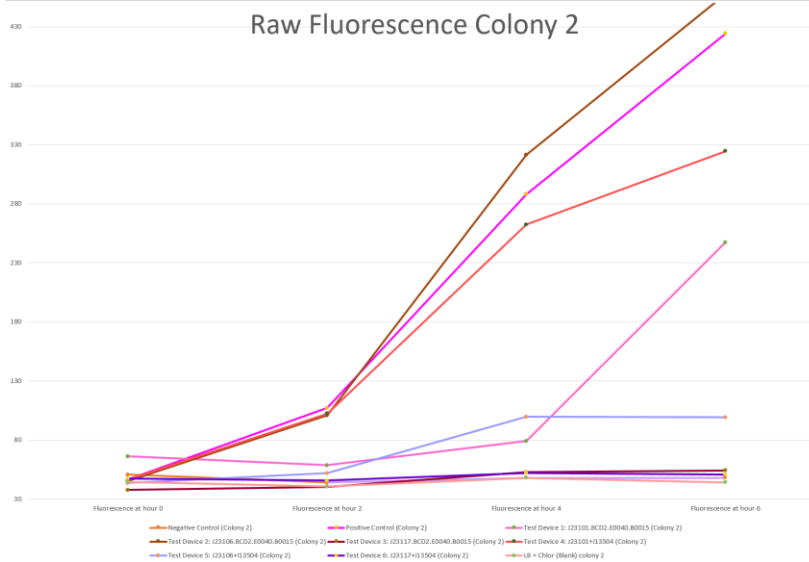
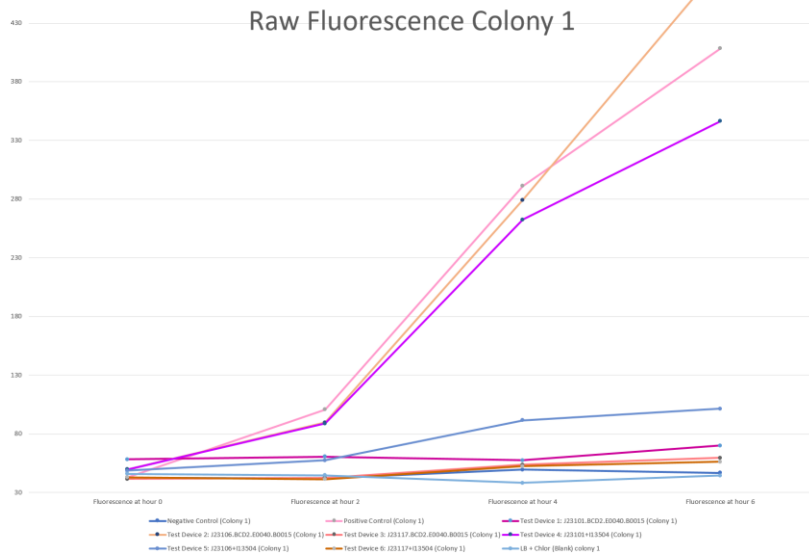
Fluorescence and OD600 in 96 holes plate

Negative control colony 1 replica 1	Positive control colony1 replica1	Device 1 colony1 Replica1	Device2 colony1 Replica1	Device3 colony1 Replica1	Device4 colony1 Replica1	Device5 colony1 Replica1	Device6 colony1 Replica1	LB +Chlor Blank colony1 Replica1
Negative control colony 1 replica 2	Positive control colony 1 replica 2	Device 1 colony 1 replica 2	Device2 colony 1 replica 2	Device3 colony 1 replica 2	Device4 colony 1 replica 2	Device5 colony 1 replica 2	Device6 colony 1 replica 2	LB +Chlor Blank colony 1 replica 2
Negative control colony 1 replica 3	Positive control colony 1 replica 3	Device 1 colony 1 replica 3	Device2 colony 1 replica 3	Device3 colony 1 replica 3	Device4 colony 1 replica 3	Device5 colony 1 replica 3	Device6 colony 1 replica 3	LB +Chlor Blank colony 1 replica 3
Negative control colony 1 replica 4	Positive control colony 1 replica 4	Device 1 colony 1 replica 4	Device2 colony 1 replica 4	Device3 colony 1 replica 4	Device4 colony 1 replica 4	Device5 colony 1 replica 4	Device6 colony 1 replica 4	LB +Chlor Blank colony 1 replica 4
Negative control colony 2 replica 1	Positive control colony 2 replica 1	Device 1 colony 2 replica 1	Device2 colony 2 replica 1	Device3 colony 2 replica 1	Device4 colony 2 replica 1	Device5 colony 2 replica 1	Device6 colony 2 replica 1	LB +Chlor Blank colony 2 replica 1
Negative control colony 2 replica 2	Positive control colony 2 replica 2	Device 1 colony 2 replica 2	Device2 colony 2 replica 2	Device3 colony 2 replica 2	Device4 colony 2 replica 2	Device5 colony 2 replica 2	Device6 colony 2 replica 2	LB +Chlor Blank colony 2 replica 2
Negative control colony 2 replica 3	Positive control colony 2 replica 3	Device 1 colony 2 replica 3	Device2 colony 2 replica 3	Device3 colony 2 replica 3	Device4 colony 2 replica 3	Device5 colony 2 replica 3	Device6 colony 2 replica 3	LB +Chlor Blank colony 2 replica 3

Negative control colony 2 replica 4	Positive control colony 2 replica 4	Device 1 colony 2 replica 4	Device2 colony 2 replica 4	Device3 colony 2 replica 4	Device4 colony 2 replica 4	Device5 colony 2 replica 4	Device6 colony 2 replica 4	LB +Chlor Blank colony 2 replica 4
-------------------------------------	-------------------------------------	-----------------------------	----------------------------	----------------------------	----------------------------	----------------------------	----------------------------	------------------------------------

## Results





**August.12<sup>th</sup>**

**Protein purification** :protein could not be washed and pass through the filter in the tube

**Made protein gel and gel electrophoresis**

## August.14<sup>th</sup>

### **Made protein gel and gel electrophoresis**

**Made cell culture media:** the container did not go through the sterilization, so mold grew inside the medium after one month.

- Liquid
- Solid

**Protein induction:** none of them reach the time of adding IPTG

- NT7
- CT7
- CFluc
- NFluc

## August.15<sup>th</sup>

**Protein induction** NT7 and CT7 were overtime

- NT7
- CT7
- CFluc
- NFluc

### **Practiced streak plate method**

## August.15<sup>th</sup>

### **Added IPTG**

- CFluc
- NFluc

### **Protein induction**

- NT7
- CT7



## August.18<sup>th</sup>

### Added IPTG

- NT7
- CT7

### Protein ultrasonic crushing

- CFluc
- NFluc

### Made polyacrylamide gel, gel electrophoresis, protein purification

- CFluc
- NFluc

## Sept.9<sup>th</sup>

### Enzyme digestion (30 $\mu$ l) using Eco311 (Bsa1), 8:45 -7:30

- NT7
- CT7
- CFluc
- Pet28a 1
- Pet28a 2
- Pet28a 3
- Pet28a 4

	Concentration of pcr product	Volume of pcr product ( $\mu$ l)	Volume of enzyme ( $\mu$ l)	Volume of FD buffer ( $\mu$ l)	Volume of ddH2O ( $\mu$ l)
NT7	55.6	18	1	3	8
CT7	114.9	8.7	1	3	17
NFluc	55.2	18.11	1	3	8
CFluc	90.3	11	1	3	15
pet 28a	63.5	10	1	3	16

### Concentration after enzyme digestion

	ng/ $\mu$ l	A260/A280	A260/A230
HNT7	23.3	1.93	0.75
HCT7	55.7	1.71	0.64

HNfluc	17.4	1.96	0.37
HCfluc	23.8	1.97	0.26
H pet28a vector	14.4	1.97	0.58

## Sept.10<sup>th</sup>

### DNA ligation (20µl) 13:30-17:00

	Inserted sequence (µl)	pet28a vector (µl)	T4 ligase (µl)	T4 ligase buffer (µl)	ddH2O (µl)
HNT7	2.2ul	3.5ul	1	2	11.3
HCT7	0.9ul	3.5	1	2	12.6
HNfluc	2.9ul	3.5	1	2	10.6
HCfluc	2.1ul	3.5	1	2	11.4

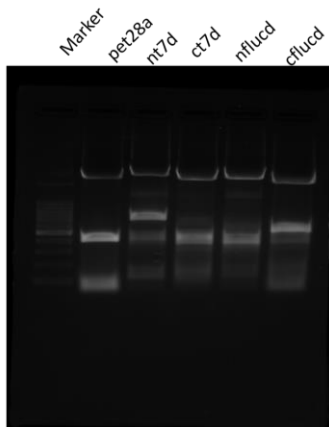
**Transformed dna** into competent cell and **incubated** around 37 celcius

- NT7
- CT7
- NFluc
- CFluc

## Sept.14<sup>th</sup>

**Transformed plasmid** into competent cell(DH5α) and **incubated** around 37 celcius (Kanamycin)

- NFluc
- dCas9
- pet281-CT7-dCas9
- control



20170914 pcr pet28a nt7d ct7d nfd cfd

**Sept.15<sup>th</sup>**

**Colony PCR (375 $\mu$ l)** NT7 grew the least bacteria, picked up 6 each of the other plates

Volume:  $\mu$ l

pcr buffer	37.5
dNTPs	30
primers	15 + 15
rTaq	3.75
ddH2O	275(273.5)

- rTaq mix: Mixed well using vortex machine and put in -20 Celsius degree refrigerator using petVF and dCas9VR

**length**

CFluc	1000bp
NT7	2250bp
CT7	1500bp
NFluc	1800bp

**gel electrophoresis** order, 7  $\mu$ l for each hole

- Marker
- CT7 1
- CT7 2
- CT7 3
- CT7 4
- CT7 5
- CT7 6
- Marker
- Nfluc 1
- Nfluc 2
- Nfluc 3
- Nfluc 4
- Nfluc 5
- Nfluc 6
- Marker
- Cfluc 1
- Cfluc 2
- Cfluc 3
- Cfluc 4
- Cfluc 5
- Cfluc 6

- Control 1
- Control 2

## **Sept.16<sup>th</sup>**

**Sway bacteria from successful pcr product from Sept.15<sup>th</sup> Kanamycin, 3 of each**

- CT71
- CT72
- CT73
- NFluc1
- NFluc2
- NFluc3
- CFluc2
- CFluc4
- CFluc6

**Redo colony PCR** control and NT7 were

- NT71
- NT72
- NT73
- NT74
- Control 1
- Control 2
- Control 3
- Control 4

**gel electrophoresis** order

- Marker
- NT7 1
- NT7 2
- NT73
- NT7 4
- Marker
- Control 1
- Control 2
- Control 3
- Control 4
- CT7
- NT7

**GFP/EFOR RED/LACZ synthesis** using Q5 polymerase, LacZ needs to go through PCR twice

## H NT7-dCas9 and H CT7-dCas9 biobrick construction

- not sure if NT7 and pet28a were connected to dCas9 or not, we tried to prove using gel electrophoresis, but without enzyme digestion, the result might be incorrect

**Gel electrophoresis (second time)** the gel was not put into the refrigerator, so it is too light

**gel electrophoresis** order; control 2 and 3 were not enough

- Marker
- CT7
- NT7
- Marker
- NT7 1
- NT7 2
- NT73
- NT7 4
- Marker
- Control 1
- Control 4

## PCR and gel electrophoresis of NT7 Biobrick/CT7 Biobrick/ LacZ/GFP/ Efored First time PCR

- GFP
- EFORED
- LacZ

**gel electrophoresis** order

- Marker
- NT7 Biobrick
- CT7 Biobrick
- LacZ
- GFP
- Efored

**Gel extraction** of CT7 biobrick

**Sept.17<sup>th</sup>**

**Plasmid extraction** plasmid were put in the box of histag

- H CFluc dCas9 1
- H CFluc dCas9 2
- H CFluc dCas9 3
- H NT7 dCas9 3
- H NT7 dCas9 4
- H NFluc dCas9 1
- H NFluc dCas9 2
- H NFluc dCas9 3
- H CT7 dCas9 1
- H CT7 dCas9 2
- H CT7 dCas9 3

**Second time PCR of LacZ** failed twice, we could not observe the result from the photo

### Sept.30<sup>th</sup>

**Transformed plasmid** into competent cell(DH5 $\alpha$ ) and **incubated** around 37 celcius;

Usage:6 $\mu$ l plasmid each, 45 $\mu$ l competent cell. Also made a control.

- NFluc dCas9
- CFluc dCas9
- NT7 dCas9
- CT7 dCas9

### Oct.1<sup>st</sup>

**Transformed plasmid** into competent cell(bl21) and **incubated** around 37 celcius;

- Pet28a-NT7-dCas9
- CT7-dCas9
- NFluc-dCas9
- CFluc-dCas9

**PCR** using Q5 polymerase

### **Enzyme digestion and gel extraction**

- va23-psb1c3
- got psb1c3 plasmid backbone

### Oct.2<sup>nd</sup>

## **Pre-culture** of bl21 bacteria

- Pet28a-NT7-dCas9
- CT7-dCas9
- NFluc-dCas9
- CFluc-dCas9

## **Gel extraction and enzyme digestion** using EcoR1 and Pst1

- Psb1c3-effored
- Psb1c3-amilgfp

## **Protein induction** IPTG 0.2 um

- IPTG 0.2 um
- NT7-dCas9
- CT7-dCas9
- NFluc-dCas9
- CFluc-dCas9

## **Transformed plasmid** into competent cell(bl21) and **incubated** around 37 celcius;

- Pet28a-shrpn-dCas9
- Pet28a-shrpc-dCas9

## **Oct.3<sup>rd</sup>**

## **Protein gel and gel electrophoresis**

- NFluc-dCas9 and CFluc-dCas9 had obvious expression

## **DNA ligation, transformed the plasmid** into competent cell and **incubated** around 37 celcius;

- Psb1c3-effored
- Psb1c3-amilgfp

## **Protein pre-culture**

- Shrpn-dCas9

## Seed-culture

- After 12hrs

## Cell-free system test (promega l1020)

- The kit did not work

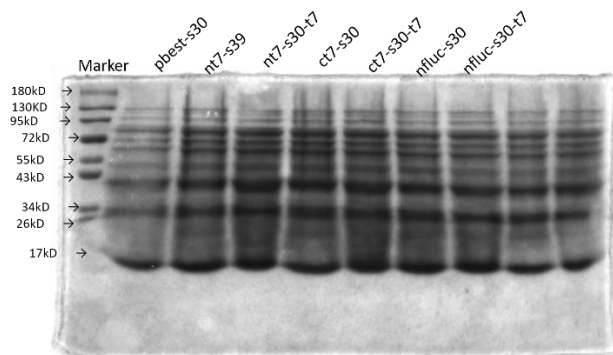
## Oct.4<sup>th</sup>

**Protein purification** using kit from takara and **buffer exchange** using 30 K MWCO concentrator from thermo

- NT7-dCas9
- CT7-dCas9
- NFluc-dCas9
- CFluc-dCas9
- The amount of protein we got were lower than 0.5ng/ $\mu$ l

## Cell-free system test

- The kit did not work again

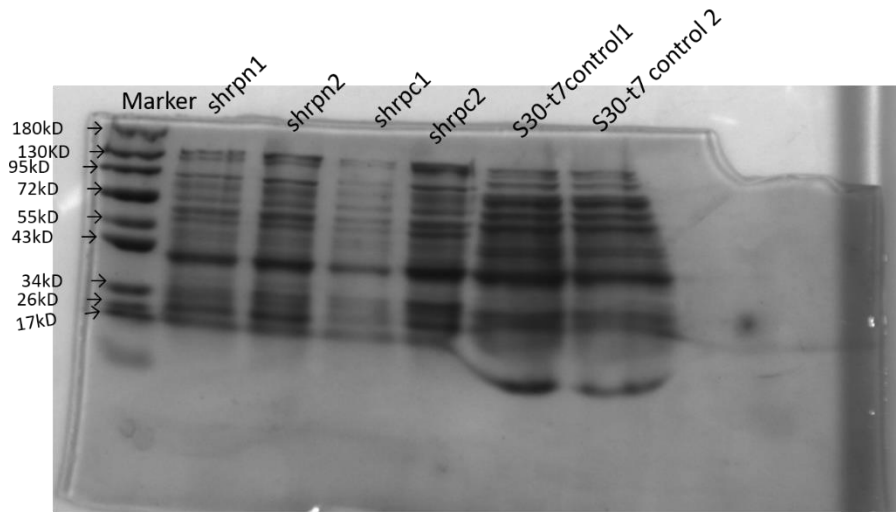


10.4 cell free M; pbest-s30; nt7-s39; nt7-s30-t7; ct7-s30; ct7-s30-t7; nfluc-s30; nfluc-s30-t7; cfluc-s30; cfluc-s30-t7

**Made protein gel and gel electrophoresis** using protein induction product

- 0.2 $\mu$ m is enough for us to induce protein





10.4 shrpm12 shrpc12 s30-t7 control12

## Colony PCR

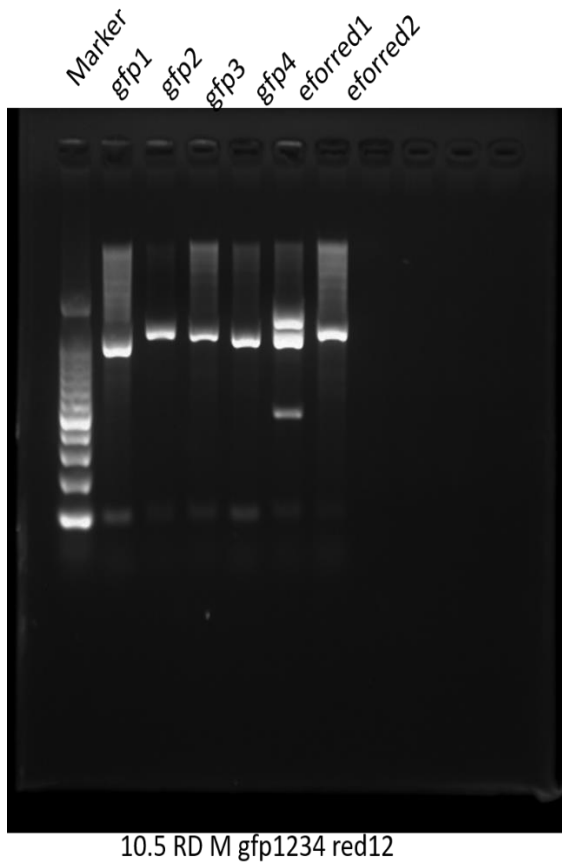
- Successfully elected correct effered plasmid

## Oct.5<sup>th</sup>

### Plasmid extraction

**Enzyme digestion** using EcoR1 and Pst1

- One of effered was conducted successfully



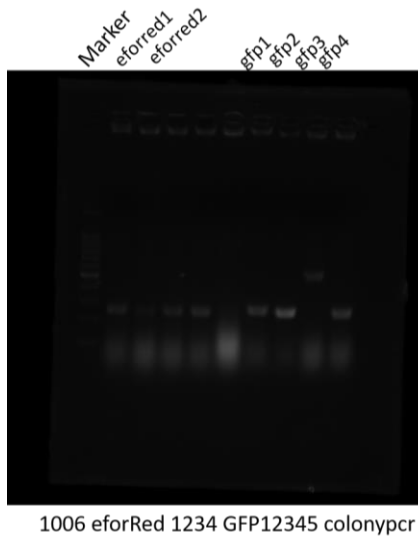
### Protein purification and gel electrophoresis of hrp

- Successful

Oct.6<sup>th</sup>

### Colony PCR

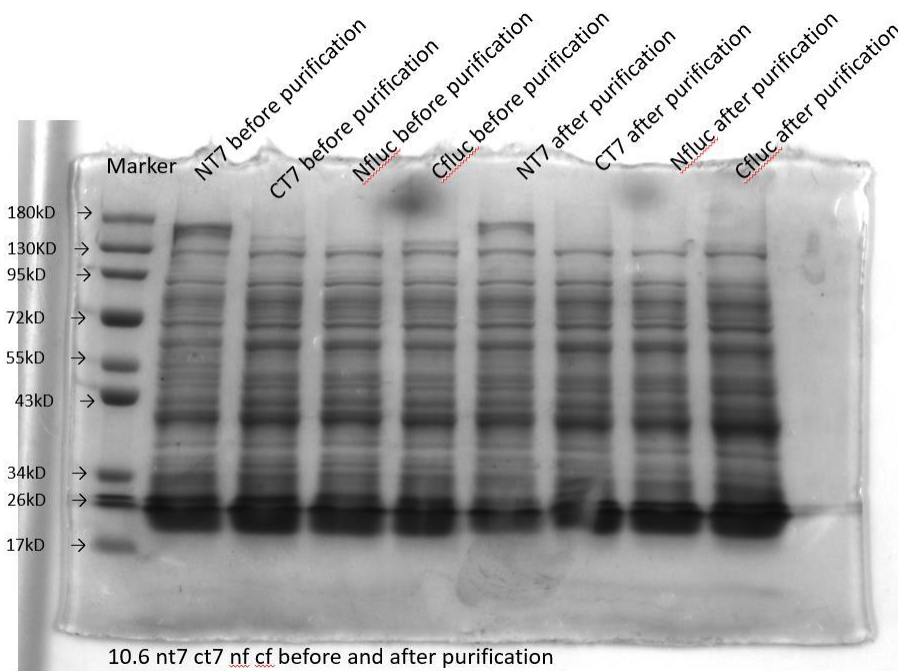
- Determined the correct colony



**Inoculation and pre-culture** bacteria from colony PCR

**Tried to find the reason of low concentration from protein purification** using protein gel before and after protein purification from sds-page

- Ability of attaching protein on the tube was really low for the takara protein purification kit. Protein could not bind to the filter membrane easily



**Protein induction**

- Pet28a-shrpn-dCas9
- Pet28a-shrpc-dCas9
- CT7-dCas9
- NFluc-dCas9
- CFluc-dCas9
- NT7-dCas9

### **Tested the effect of concentrator**

- Concentrator could not dilute protein with low molecular weight, also too much protein would block the filter membrane

### **Oct.7<sup>th</sup>**

### **Collected the correct plasmid and enzyme digestion** using EcoR1 and Pst1

- Psb1c3-amilgfp
- Psb1ce-eforred
- Successfully constructed

### **Extracted correct plasmid and sent to BGI for sequencing**