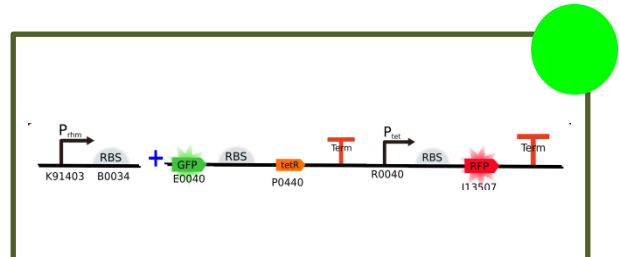


## Assembly:

Pr\_I13507



## 1<sup>st</sup> Day:

EXSP Digestion (see [Enzymatic Digestion Protocol](#))

	Part	Size	ng/μl
1	Pr_RBS	134 bp	74.6
2	GFP_I13507	2585 bp	273.5

	Volume to 1,0 μg (μl)	Buffer 10x (μl)	BSA	Enzime 1	Volume (μl)	Enzime 2	Volume (μl)	H2O to 20μl (μl)
1	13.4	2 (M)	-	SpeI	1	PstI	1	2.6
2	4	2 (M)	-	XbaI	1	PstI	1	12

Final Plasmid	Resistance
pSB1A2	ampicillin

## Gel purification

- See PureLink® Quick Gel Extraction Kit Invitrogen™ manual
- Quantify digestion products

Parts	ng/μl
Pr_RBS	5.1
GFP_I13507	12.3

**Obs:** 260/280 in a quality parameter that tells you if your sample is contaminated with proteins. The greater it is compared to 1 the less contaminants you have.

## Ligation (see Ligation Protocol)

Part containing the plasmid	Pr_RBS	9.5 $\mu$ l
Insert	GFP_I13507	6 $\mu$ l
10x T4 DNA Buffer		4 $\mu$ l
T4 DNA ligase 1u		0.5 $\mu$ l
H2O to 20 $\mu$ l		-

**Obs:** To determinate the amount of DNA necessary we used the following equation

$$\text{Insert ng} = \text{plasmid ng} \times \text{insert bp} / \text{plasmid bp} \times \text{insert: plasmid ratio}$$

- Incubate overnight at 37°C.
- Prepare and sterilize in the autoclave tubes with 6 ml of liquid LB medium.
- Prepare glycerol 40%

## 2<sup>nd</sup> Day:

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Transformation (see Transformation Protocol in Escherichia coli DH5- $\alpha$ )

- Organism: E. coli DH5- $\alpha$
- Selection: Ampcillin

## 4<sup>th</sup> Day:

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Confirmation with NotI