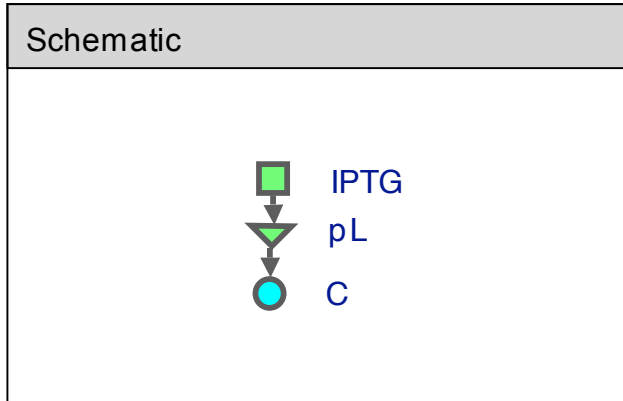


[Trc-LC] Inducible CFP Expression Device

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	Host: Ecoli K12Z1
	p1.lacI::p2.tetR
	Part: [Trc-LC]
	pL.C
	Small molecules
	<ul style="list-style-type: none"> •IPTG: Isopropyl β-D-1-thiogalactopyranoside.
	Promoters
	<ul style="list-style-type: none"> •p1: Constitutive LacIq promoter. •p2: Constitutive N25 promoter. •pL: Lac promoter.
	Proteins
	<ul style="list-style-type: none"> •LacI: Lac repressor, negative regulator of pL. •TetR: Tet repressor, negative regulator of pT. •C: Cyan fluorescent protein
	Description
	IPTG drives the expression of CFP.
	Usage and compatibility
	This device can be used to calibrate fluorescence readings.
<p>Registry ID: BBa_I726005</p>	

Characteristics

Protocol: We grew cells overnight in LB. We then transferred them to Glu-M9 minimal medium containing the desired final concentration of IPTG, and allowed them to grow for 12h. The cell density at transfer was chosen so that the final OD600 was < 0.1. Cells were concentrated by centrifugation, and imaged on an epifluorescence microscope. We calculated the fluorescence per unit area of single cells, obtaining data from ~500 cells for each run. We then averaged these values in log space to obtain the final estimate of protein expression. The IPTG mesh was [0 5 10 50 500 1000] uM.

Sigmoidal fit:

$$y = a_0 + a_1 \frac{x^n}{K^n + x^n} \quad a_0 = 96; a_1 = 818; K = 158; n = 2;$$

Measurements and analysis were carried out by members of the NCBS iGEM 2007 team:
 Kiran, Krishna, Mukund, Navneet, Nilesch, Senthil, Shashanka, Sugat, Sushant, Varun, Vini, Vivek