

iGEM 2010 Progress Report

June 21, 2010





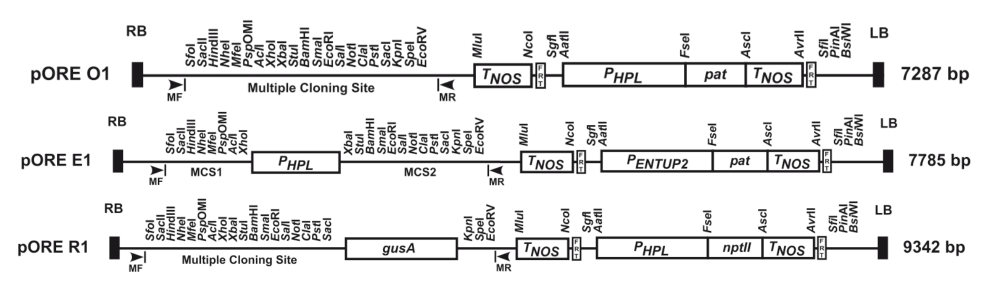






Aim: biobrick agrobacterium vector

We are working with 6 different plasmids known as pORE:

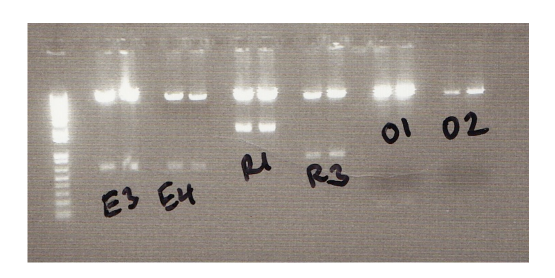


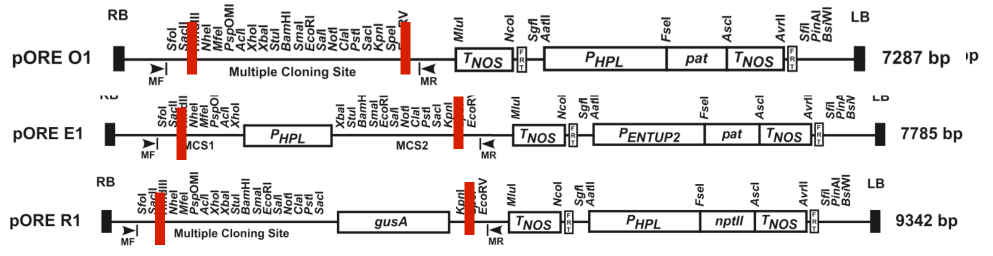
To achieve our aim we need to remove the MCS and replace it with the biobrick cloning sites

Progress so far: Backbones

Used restriction digests to remove MCS + promoters/ reporters from each plasmid, leaving the backbone.

- Open Series: cut with SacII and SpeI
- Reporter Series: cut with HindIII and Spel
- Expression Series: cut with HindIII and Spel

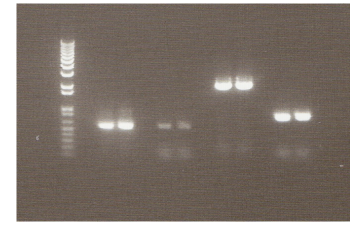




Progress: Inserts for reporter and expression series
Created inserts from primers via PCR on the original
agrobacterium vector designed to retain any reporter or
promoter already in the vector and add the Biobrick Multiple
Cloning Site

Reporter Series:

- forward: HindIII cut site, Biobrick
 MCS, first part of appropriate reporter
- reverse: last part of reporter, Nhe cut site



E3 E4 R1 R3

Expression Series:

- o forward: HindIII cut site, first part of promoter
- o reverse: last part of promoter, Biobrick MCS, Nhe cut site

Progress so far: Ligation of inserts and backbones

- Ligated inserts and backbones of expression and reporter series vectors to create biobrick agrobacterium plasmids.
- Transformed into NEB Turbo Competent E. coli cells.
- Picked colonies and made cultures.

Next steps:

- Create inserts for O1 and O2 by annealing oligos and digest with SacII and NheI.
- Ligate O1 and O2 backbones and inserts.
- Transform resulting O1 and O2 plasmids into E. coli.
- Transform each plasmid into agrobacteria.
- Sequence constructs to check for accuracy

Next: Team Flavor

TEAM FLAVOR

Our Plan:

- Express different flavors in plants
 - Targeting and altering specific metabolic pathways
 - Producing proteins
- Make BioBrick Parts

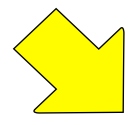
Flavors:

- Miraculin
 - Turns 'Sour' to 'Sweet'
 - Ordered from Mr. Gene
 - Arrival Pending... (should be early this week)
- Brazzein
 - 500x to 2000x sweeter than sugar (Wikipedia)
 - o Ordered from Mr. Gene
 - Arrival Pending... (should be early this week)
- Valencene
 - Orange/Citrus smell, derived from Valencia oranges
 - Extracted from oranges ourselves
- Wintergreen Scent
 - From the 2006 MIT iGEM team
 - Full pathway available in 2010 iGEM kit
- Banana Scent
 - From the 2006 MIT iGEM team
 - Parts of pathway available in 2010 kit

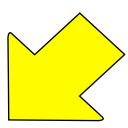
To Do:

Make BioBrick parts out of all enzymes/proteins in the flavour pathways.

Make BioBrick parts out of pORE vector components



Insert the newly created flavour BioBricks into the optimized Arabidopsis vector (courtesy of Team Vector)



Express in plants!

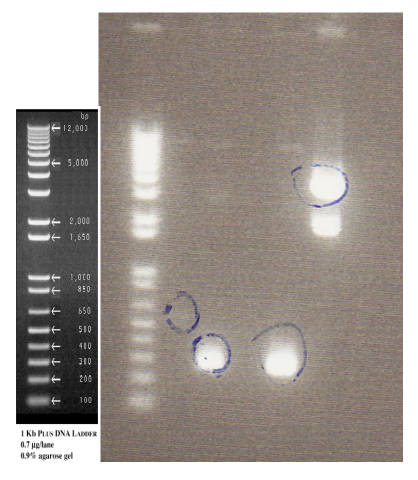
What we have done this week:

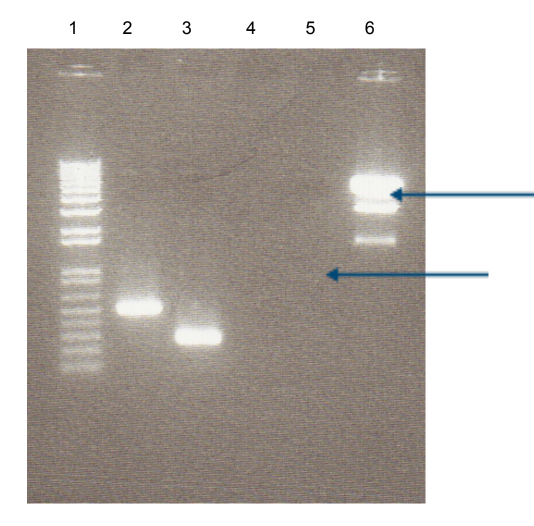
- 1. BioBrick parts from the registry were transformed and expressed in E. Coli:
 - Full and partial wintergreen pathway
 - Very few colonies
 - Parts of banana pathway
 - Expressed components of banana pathway did not grow!
 - PCR of restriction digest yielded products of incorrect size
- 2. Primers were designed, and PCR preformed on components of the e3 pORE vector
 - Promoter
 - Terminator sequence
 - Terminator sequence + Stop codon
 - We are in the process of inserting these components into standard BioBrick vectors (V0120). First attempt at ligating into vector did not work. Result of second ligation attempt will be seen on Monday.

Digestion of PCR Products

- 1. Ladder
- 2. Promoter pENTCUP2
- 3. NOSt
- 4. empty
- 5. NOSt + STOP
- 6. Plasmid

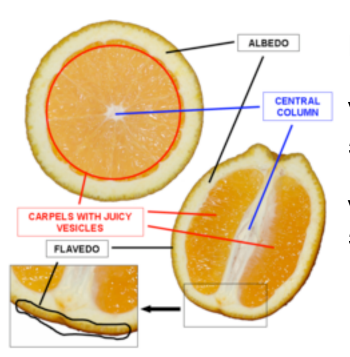
Ladder 1 2 3 4 5 6





What we have done this week (cont'd):

- 3. Valencia orange RNA extraction and cDNA synthesis
 - extracted RNA from flavedo of Valencia orange
 - need to remove pstl sites from valencene gene without altering functionality
 - site directed mutagenesis
 - primers have been ordered

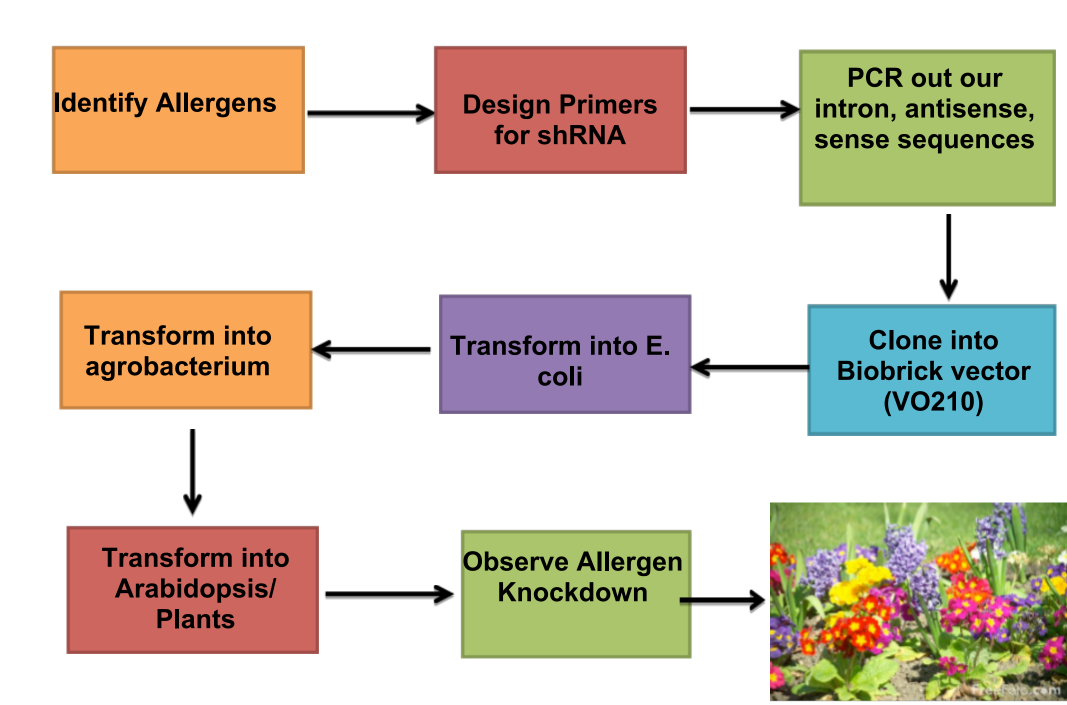


Primers for site directed mutagenesis:

Valencene synthase t30a (amino acid 440 lle -> lle) 5' gagaaacatttcgtcctacagcagatttccatcctagtt 3'

Valencene synthase c1320a (amino acid 10 Thr -> Thr)
5' aggttgtaaaagcagcatcagttatatgcagactcatgga 3'

Team Allergy



Team Allergy Goals

Short Term

Design hp RNA to target GFP; introduce to Arabidopsis; observe GFP knockdown through qrt pcr/ fluoresence

Design hp RNA to target Fra a1/LTP1; introduce to strawberry/Aribidopsis; observe knockdown through assays Long Term (allergen knockdown)

> Knockdown of GFP in GFP Arabidopsis through hp RNA interference

> Knockdown of Fra a1 (strawberry allergen) & LTP (lipid transfer protein) in Arabidopsis

What We Did...

Monday and Tuesday:

RNA extraction on strawberry fruit obtained from Shaws Obtained sufficient mRNA to yield the following amounts of cDNA



Sample #	cDNA concentration (ng/uL)	mRNA concentration (ng/uL)	Part of the strawberry
1	441	11	Outer Receptacle
2	1477	36	Receptacle
3	2749	6.5	Inner Receptacle

PCR

Wednesday and Thursday:

Tried to PCR out Fraa 1 for biobrick cloning using cDNA from strawberries

Goal: 300bp DNA sequence (Fraa1)

Result?

Unsuccessful --- Primer Dimers? PCR products were ~ 100 bp

What went wrong? We eliminated the following possibilities through further testing

- Incorrect annealing temperature by using a temperature gradient
- Too much cDNA in the reaction by reducing the amount of cDNA from micrograms to ~100 nanograms/ 50 uL reaction



One Important Question!

Was there mRNA coding for Fraa1 in the strawberries when we extracted mRNA?

Further research on Fraa1 reveals that Fraa1 is part of the ripening mechanism of strawberries

Strawberries are non-climactic fruit, which stop all ripening metabolic processes after removal from plant. Our strawberries were store-bought, and therefore have been "dead" for at least two or three days, giving the mRNA coding for Fraa1 plenty of time to degrade.

Next experiment ----

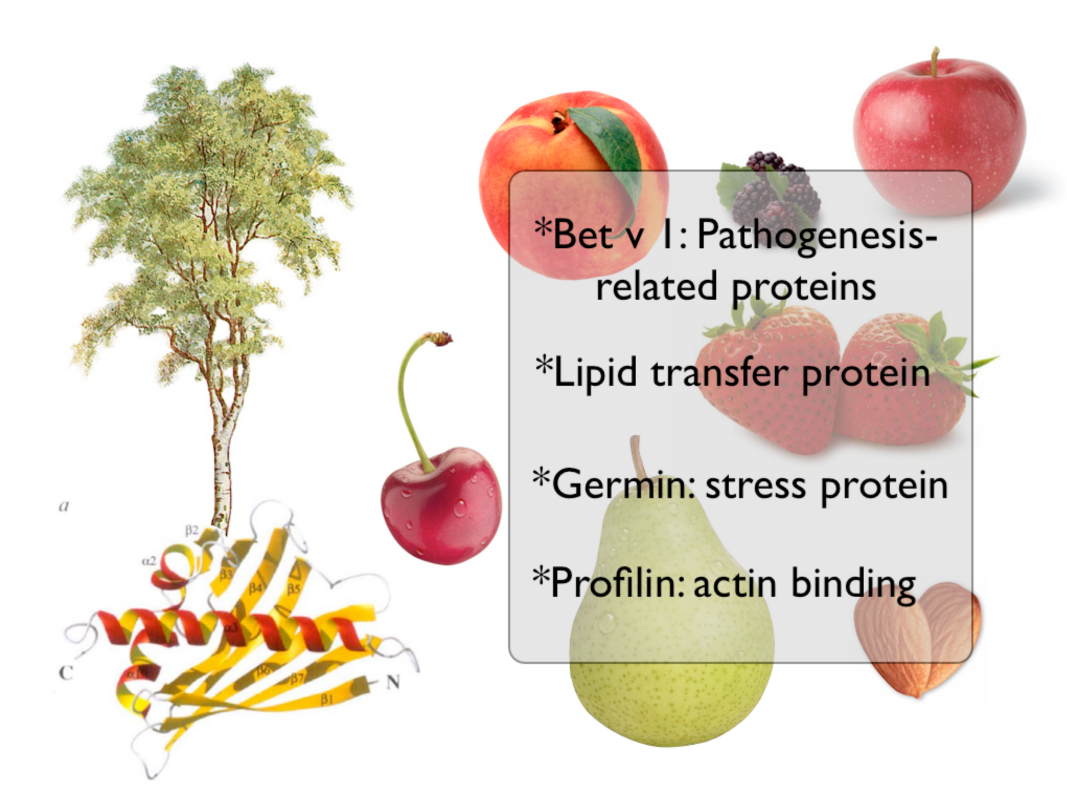
- Make sure that we have reason to believe that our sample is in a state that contains the appropriate mRNA

Onward!

Search for allergens in arabidopsis It should be ready(er) by now.

Namely:

LTP1 (documented allergen)
Bet v 1 (thaliana <= latera <= birch homologue via BLAST)
Others (list, links on wiki)













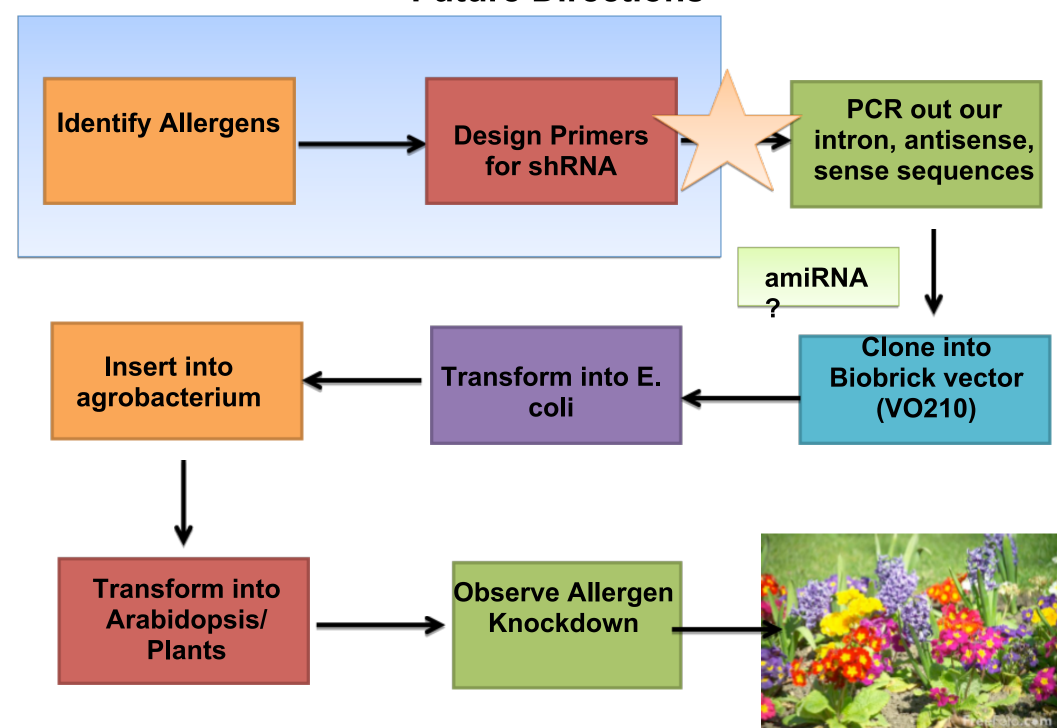


They should be larger by now.

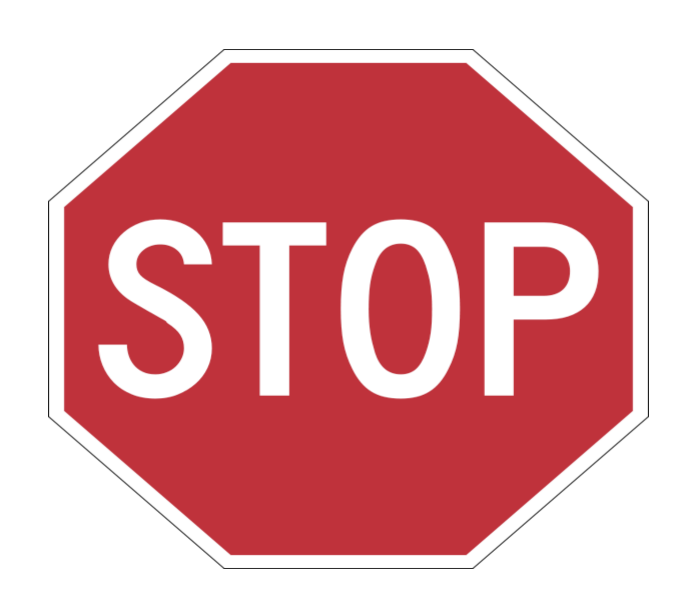
After all, they spent the weekend happily growing in the underground greenhouse bunker.

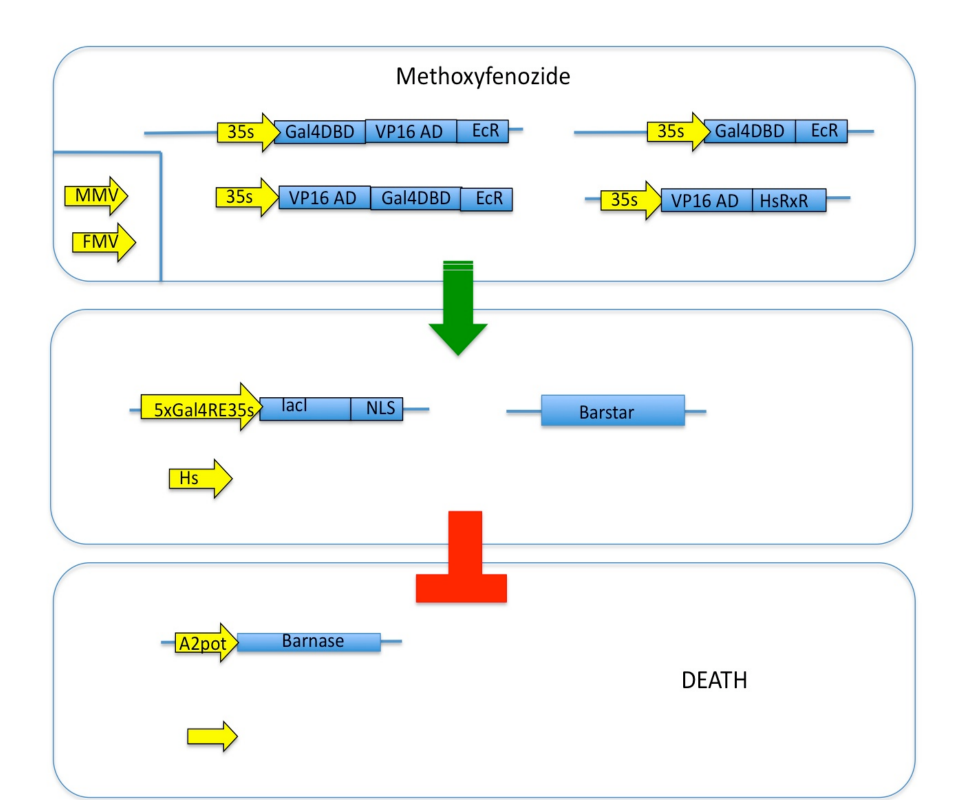


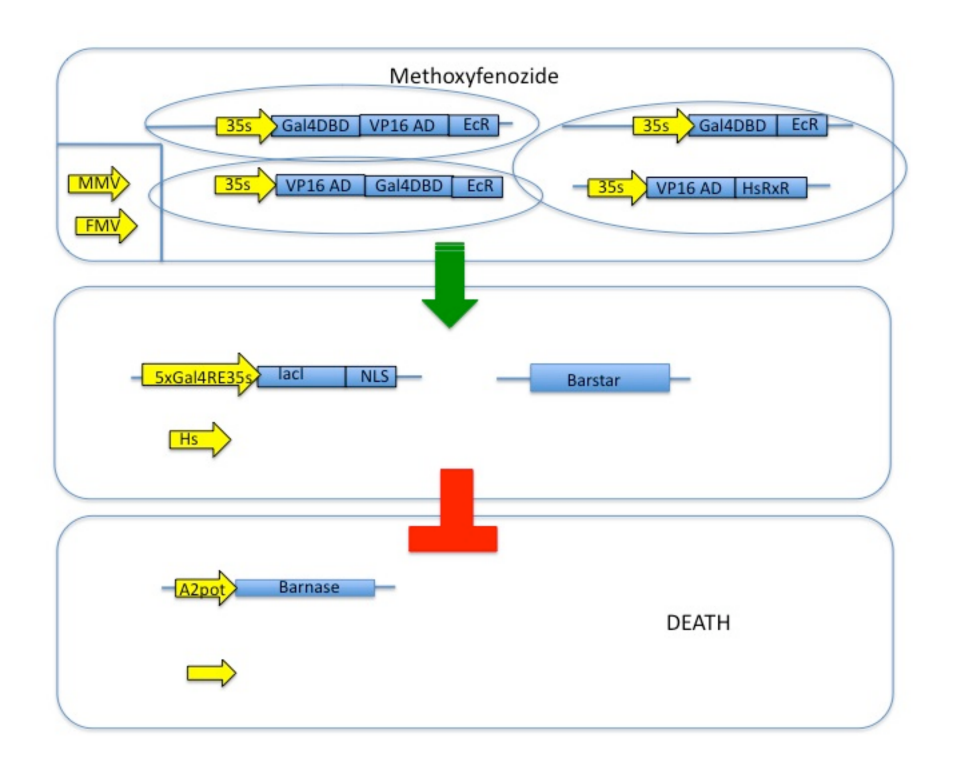
Future Directions

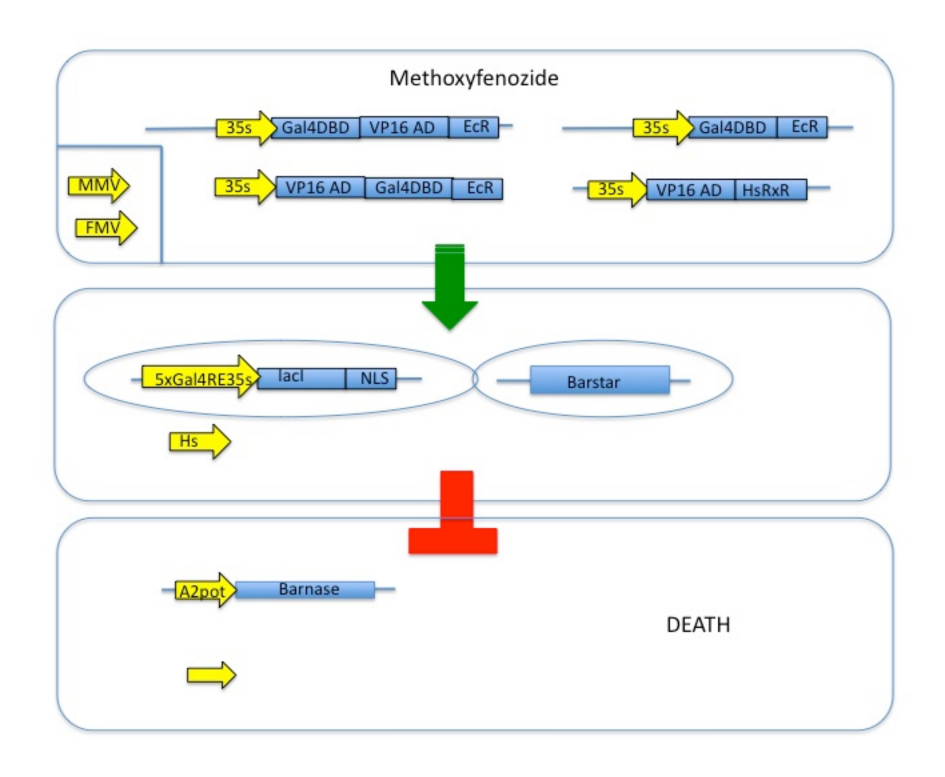


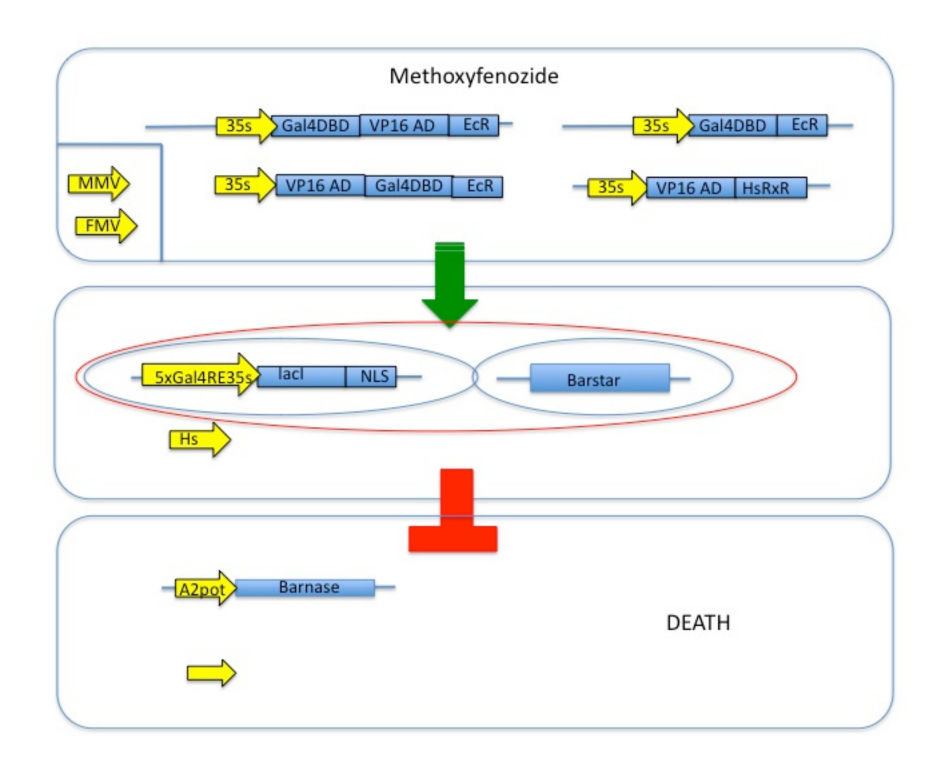
TEAM GENETIC FENCE











Last Week

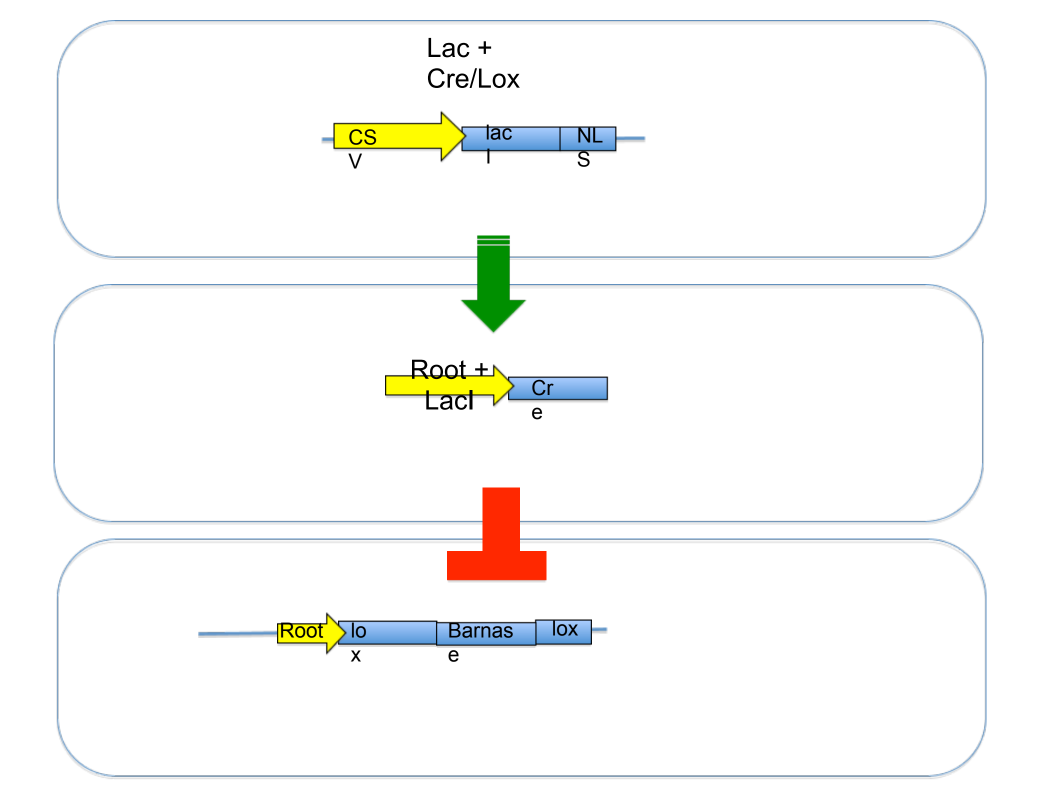
- Methoxyfenozide was ordered and arrived Thursday
- Made requests to other labs for LacI with NLS, ActIn promoter with lacO sites (A2pot), and Cassava Vein Mosaic Virus promoter, as well as plasmids containing the entire monopartate and bipartate switch mechanisms
- Barstar and Barnase ordered from ADDGENE

Last Week, Cont.

- VP16, GAL4 DNA Binding Domain, and Lacl in biobricks registry
- Lacl
 - Two biobricks in registry, one wildtype and one including and LVA rapid degradation tail
 - Transformed, minipreped and digested both Lacl and LacI+LVA
- VP16
 - Transformed, minipreped and digested VP16
- GAL4 DBD
 - Transformed
- One run of PCR to attach NLS to Lacl
 - Also annealed oligos to form NLS alone for biobrick

Alternate/Secondary System: Crelox and IPTG

- Attractive paradigm (lactose analog instead of pesticide)
- However untested and relies on less-characterized parts



TECHNICAL ADVANCE

External control of transgene expression in tobacco plastids using the bacterial *lac* repressor

Stefan K. Mühlbauer and Hans-Ulrich Koop*

Icon Genetics AG, Research Centre Freising, Lise-Meitner-Str. 30, 85354 Freising, Germany

Received 17 March 2005; revised 9 June 2005; accepted 17 June 2005.

*For correspondence (fax +49 89 17861165; e-mail koop@lrz.uni-muenchen.de).

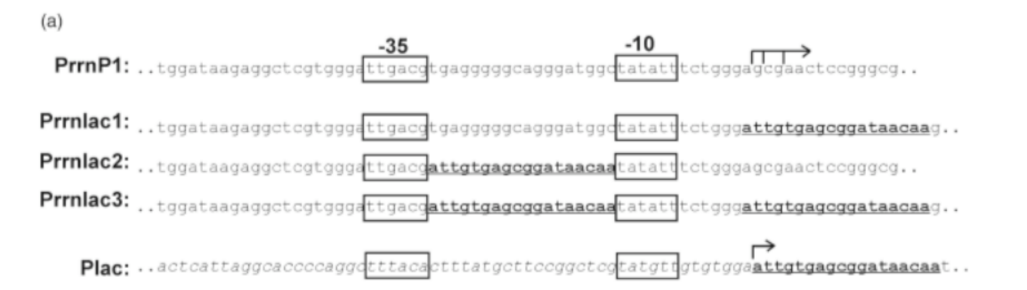
Summary

Although several induction systems have been described for plants containing transgenes in the nucleus, to date there is only one method for controlling transgene expression in plastids. This consists of chemical induction of a nuclear gene and import of the gene product into plastids, so that transformation of two cellular compartments is required. Here we describe a system for external control of plastid gene expression which is based entirely on plastid components and can therefore be established in a single transformation step. Our system uses modified promoters containing binding sites for the bacterial *lac* repressor. Chemical induction can be made with intact plants or after harvesting, which provides ecological and economic benefits.

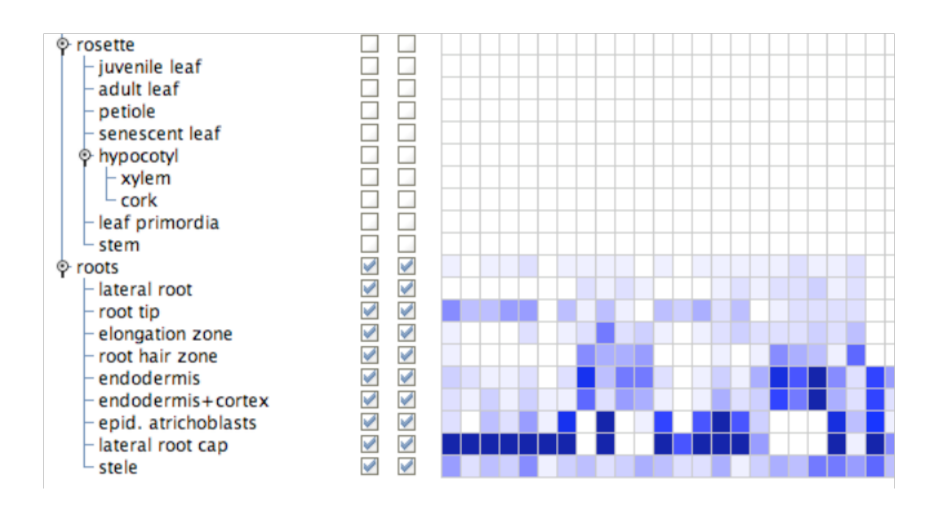
Keywords: plastid transformation, transgene expression, induction, lac repressor, IPTG.



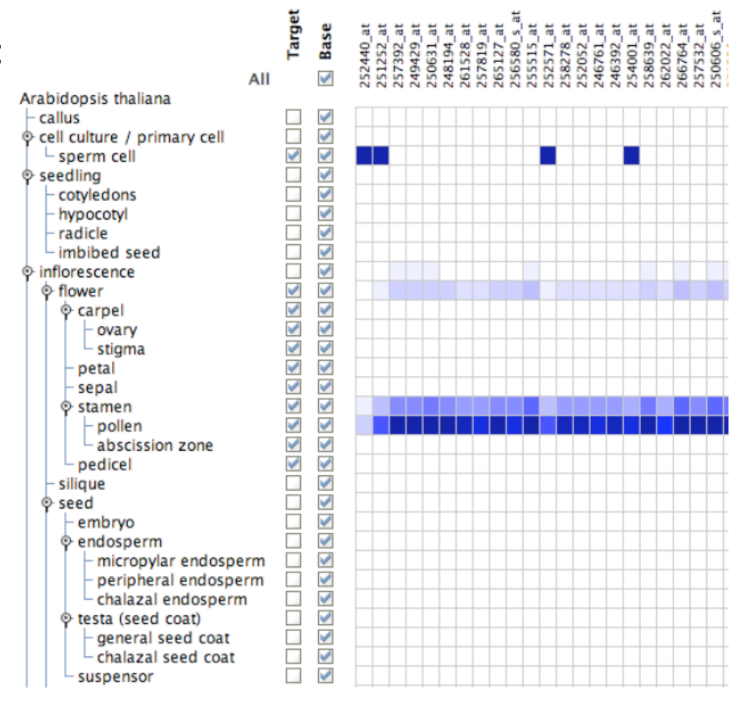
Control of plastid gene expression 943



Tissue Specific Promoters



Back up:



Other considerations

- Plastid DNA and Chloroplast modification
 - dramatically reduces risk of cross-pollenation
 - Increases technical difficulty of transformation (agrobacterial vector no longer sufficient

Time specific tissue-specific promoters for cre-lox?