

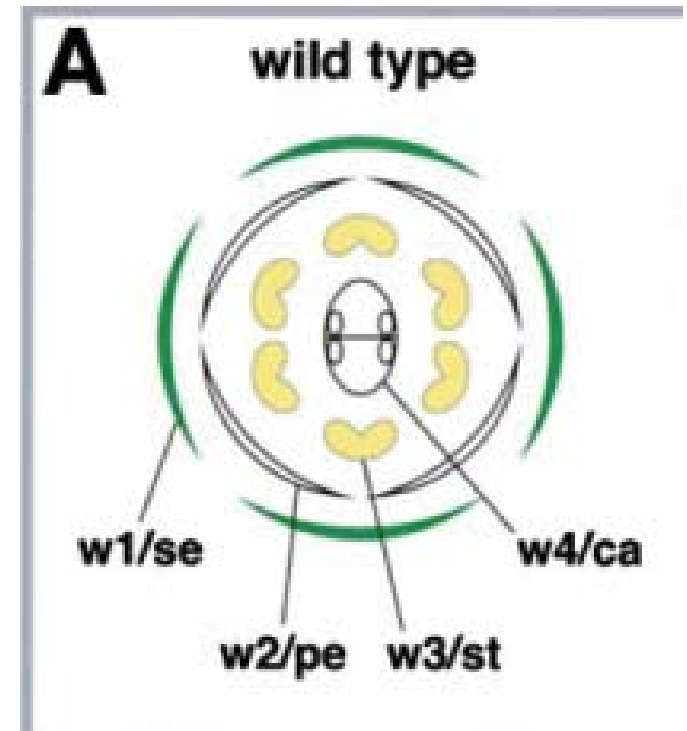
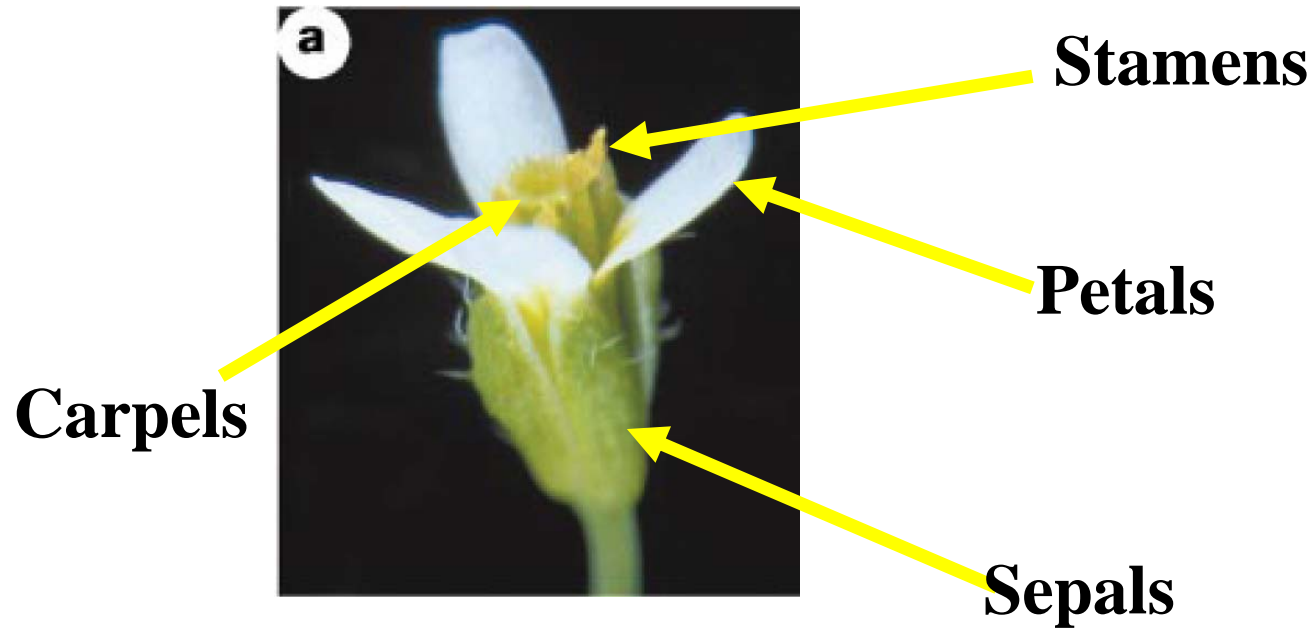


**Genetic Specification of floral organ identity**

**Initiating floral development**

**Deciding when to initiate flowering - induced mutations**  
**- in Nature**

# Flower structure of Arabidopsis



**In Arabidopsis:**

<b>Whorl 1</b>	<b>4 sepals</b>
<b>Whorl 2</b>	<b>4 petals</b>
<b>Whorl 3</b>	<b>6 stamens</b>
<b>Whorl 4</b>	<b>2 carpels</b>

# **Mutations that affect flower development placed in classes**

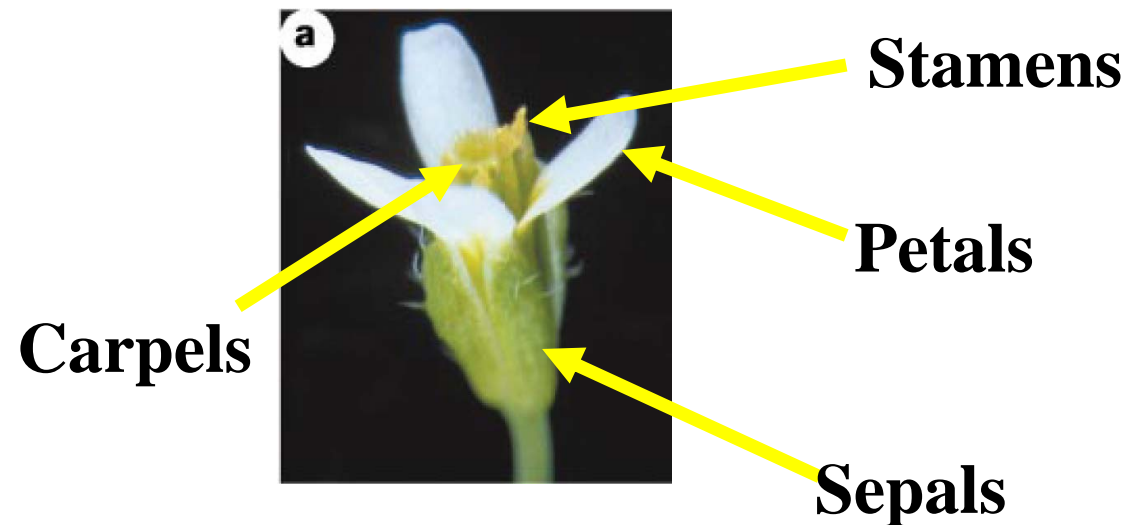
**Alter the organs formed in two adjacent whorls**

**A mutants : affect sepals and petals**

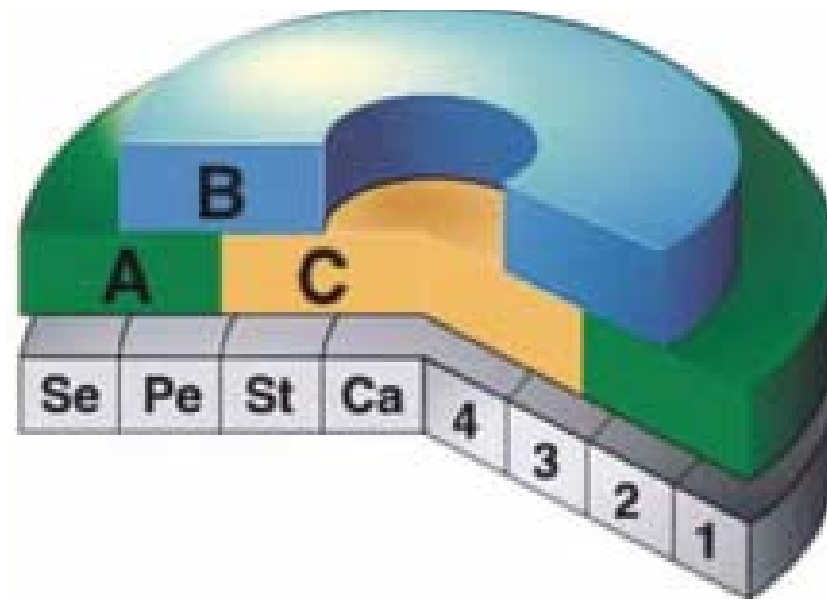
**B mutants : affect petals and stamens**

**C mutants : affect stamens and carpels**

**Antirrhinum**

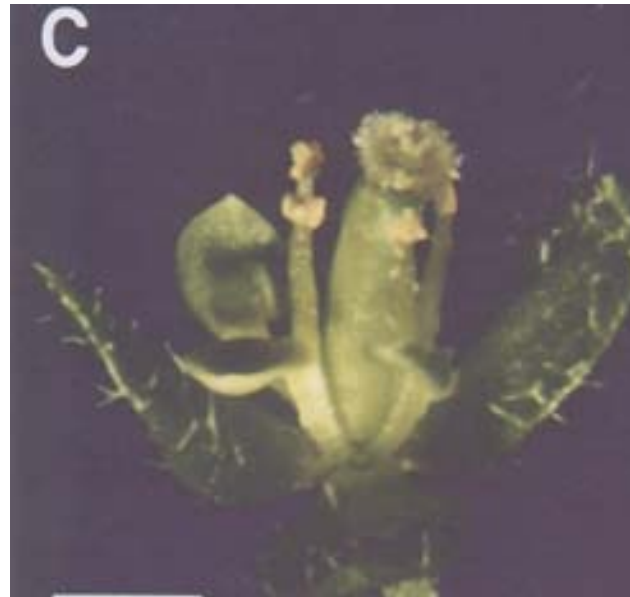
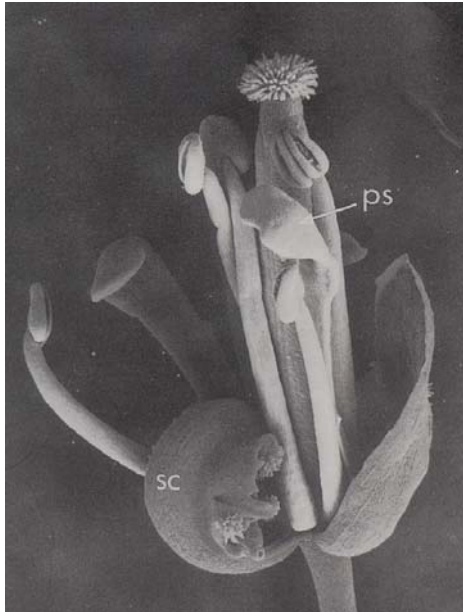


Specification of floral organ identity occurs in four circular domains called whorls and is controlled by three genetic functions called A, B and C



# A function mutations affect flower development in Whorl 1 and 2

apetala2



apetala2



Wild-type

Sepals	————→	carpelloid organs
petals	————→	Absent or stamenoid
stamens	————→	normal
carpels	————→	normal

# B function mutations affect flower development in Whorl 2 and 3

apetala3



pistillata



Wild-type

Sepals —————> normal  
petals —————> Sepals or absent  
stamens —————> carpelloid  
carpels —————> normal

# C function mutations affect flower development in Whorls 3 and 4



agamous

Sepals —————> normal

petals —————> normal

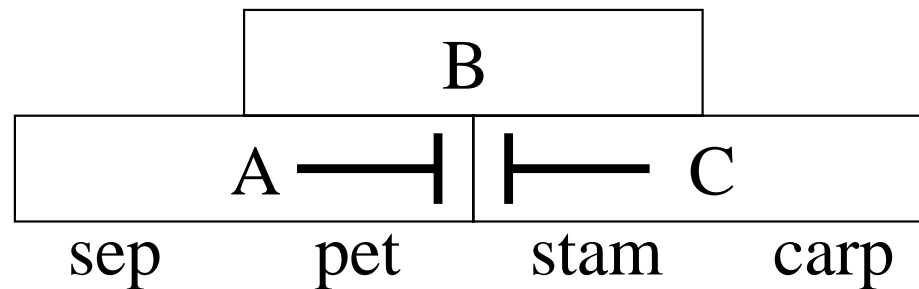
stamens —————> petals

carpels —————> Sepals and flower  
inside a flower



Wild-type

# Analysis of mutant phenotypes suggested ABC model



A function - APETALA1, APETALA2

B function - APETALA3 PISTILLATA

C function - AGAMOUS

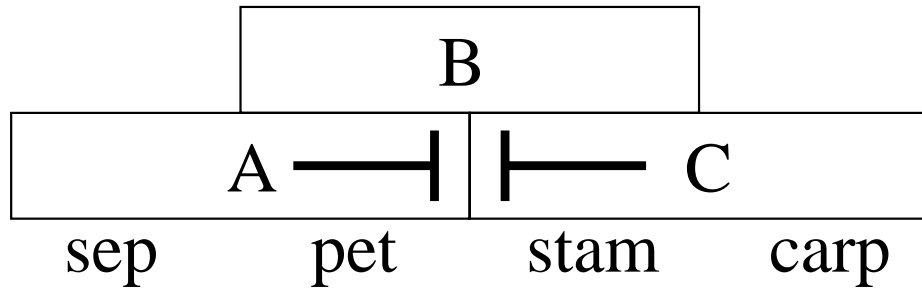
Overlapping functions provide new identities

A opposes C and C opposes A

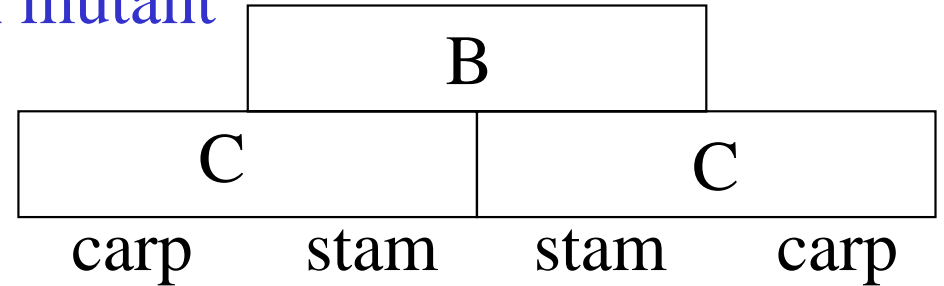


# Interpretation of mutant phenotypes according to ABC model

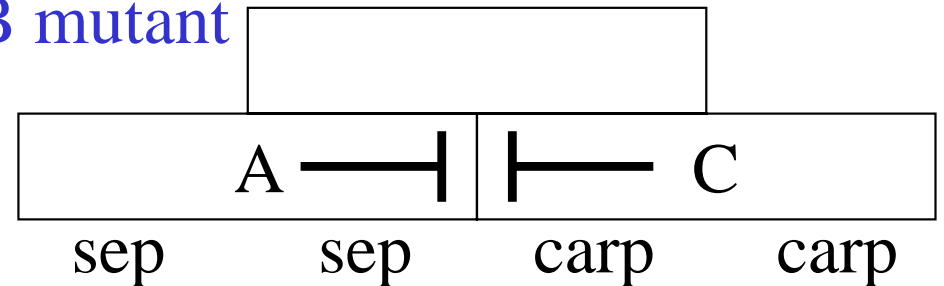
Wild type



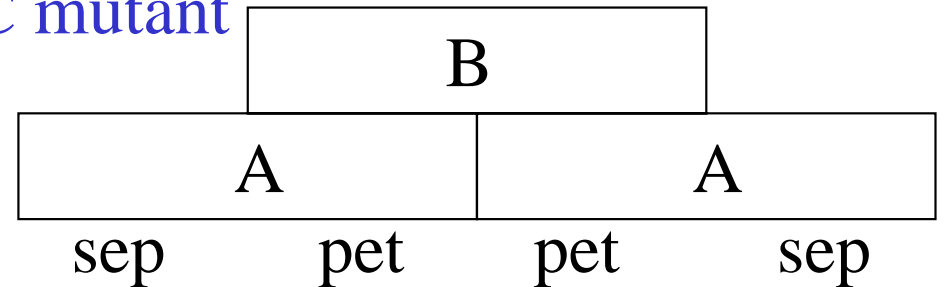
A mutant



B mutant



C mutant



# Double mutants



**pi ag double mutant / loss of B and C function**  
**All whorls are sepals**  
**More whorls**



**ap2 pi double mutant /**  
**loss of A and B function**

**All whorls are carpels**



**Wild-type**

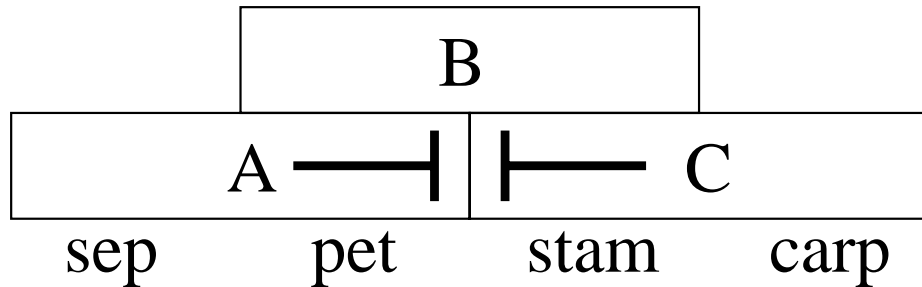


**ap2 ag double mutant / loss of A and C**

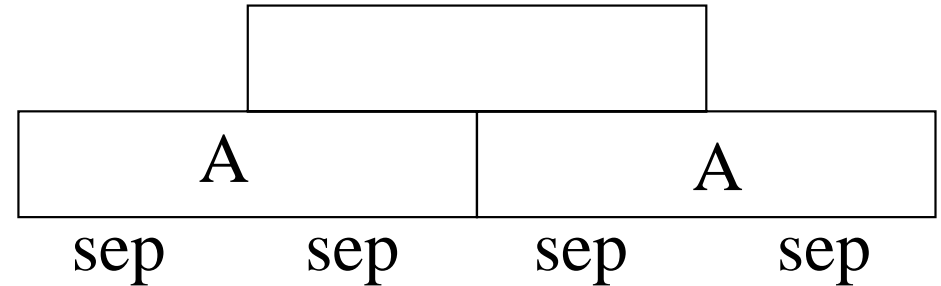
**Leaves in 1st and 4th whorl**  
**Modified petals/stamens in second and third**  
**More whorls**

# Interpretation of double mutant phenotypes according to ABC model

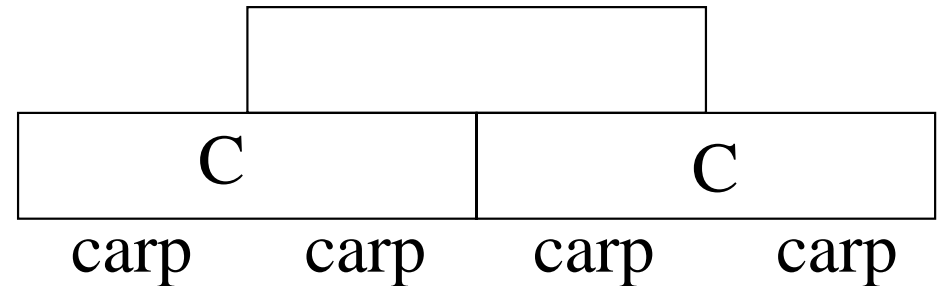
Wild type



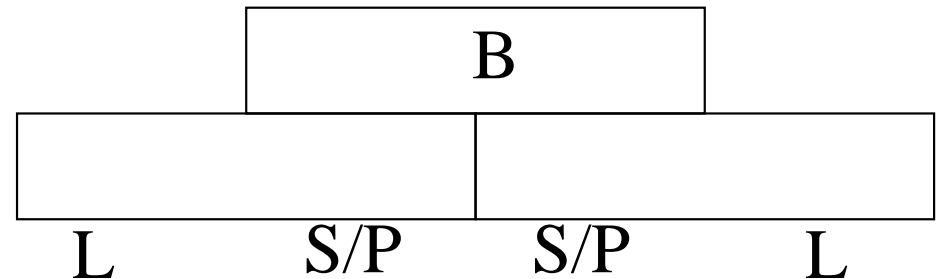
BC mutant



AB mutant



AC mutant



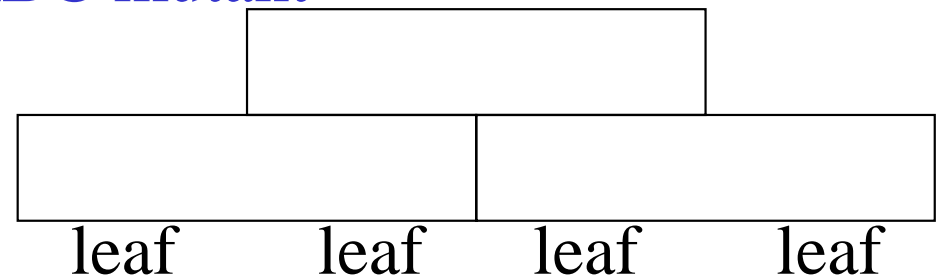
# ABC triple mutant phenotype



ap2 pi ag triple mutant / loss of ABC function

All whorls leaf like

ABC mutant



# Structure of plant MADS box transcription factors

MADS	I	K	C
------	---	---	---

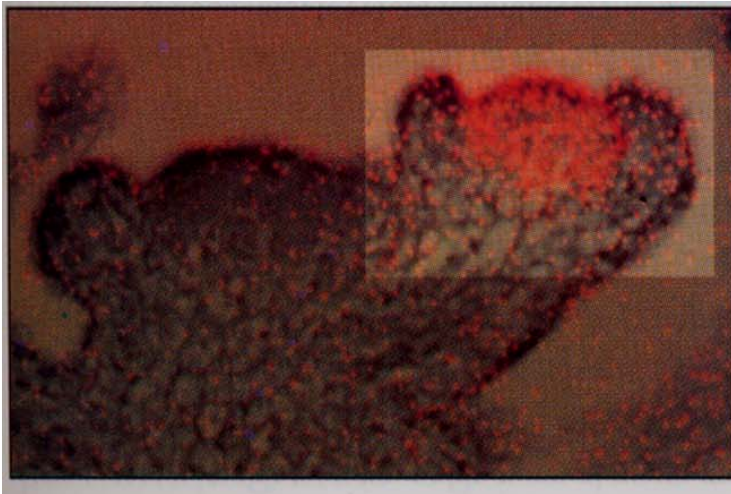
MADS transcription factor

MADS	-	DNA binding
I	-	intervening region
K	-	protein-protein interactions
C	-	carboxy-terminal domain

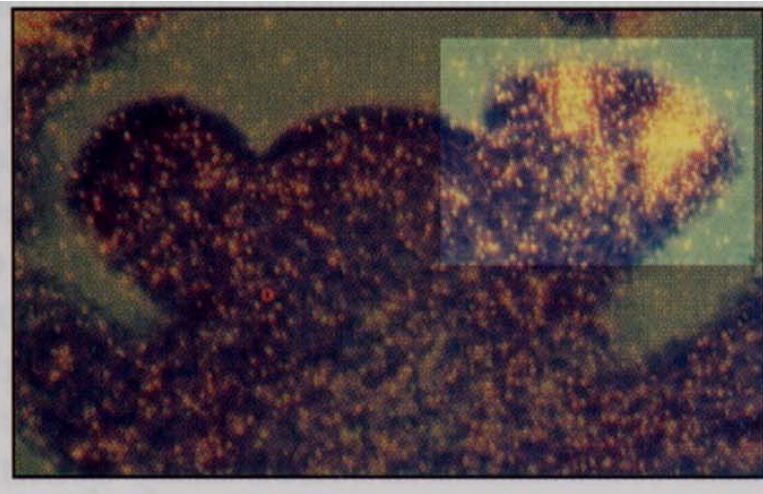
Bind DNA through a conserved DNA sequence called CArG  
Box – CC(A/T)<sub>6</sub>GG

AP1, AP3, PI, AG are MADS box transcription factors

Patterns of expression indicate that spatial control  
is regulated mainly at the level of transcription



Agamous  
C function  
Expressed in whorls 3 and 4



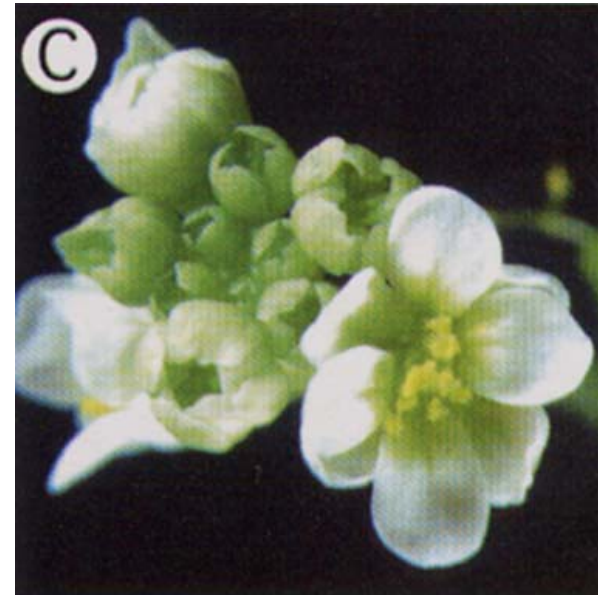
Apetala3  
B function  
Expressed in whorls 2 and 3

# Misexpression from the 35S viral promoter in transgenic plants supports proposed roles of B function genes

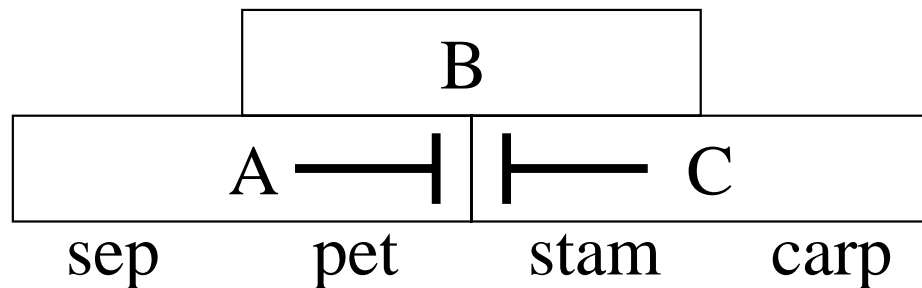
Wild type



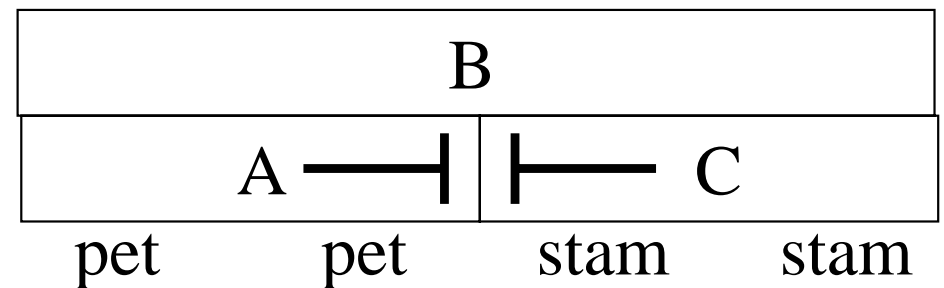
35S::AP3 35S::PI



Wild type



35S::AP3 35S::PI

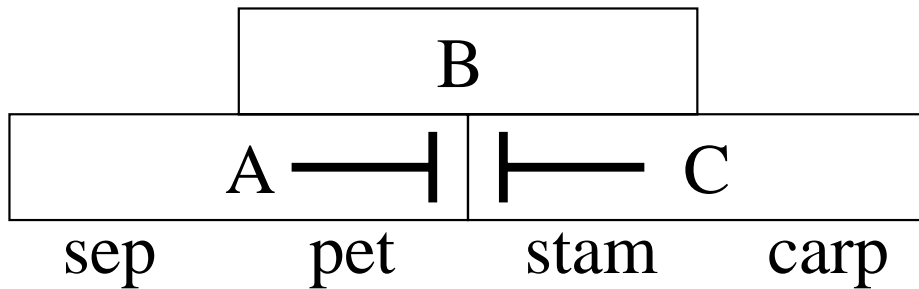




# Regulation of Floral organ identity gene expression

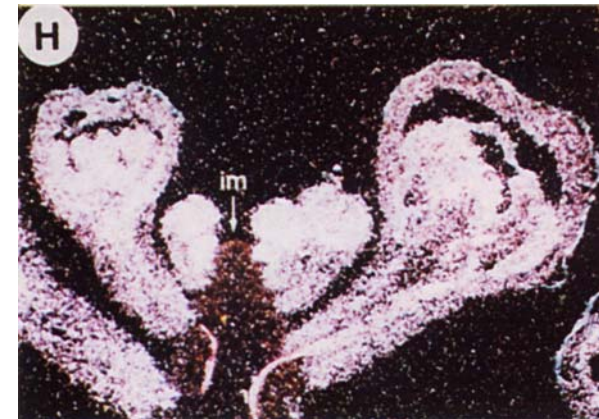
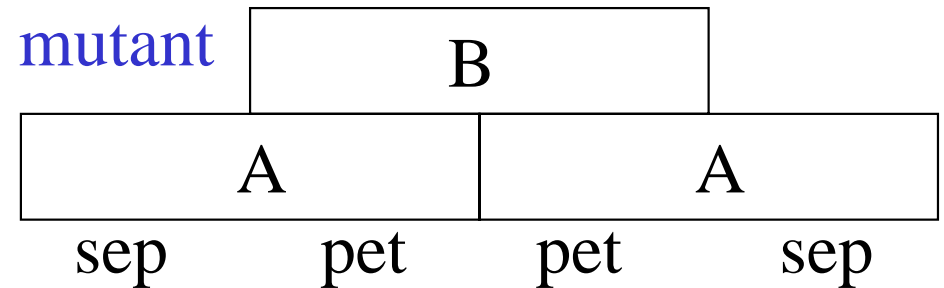
AP1 (A function) misexpression in ag (C function) mutant

Wild type



AP1 mRNA in  
Wild-type plants  
Only in whorl 1 and 2

ag  
C mutant



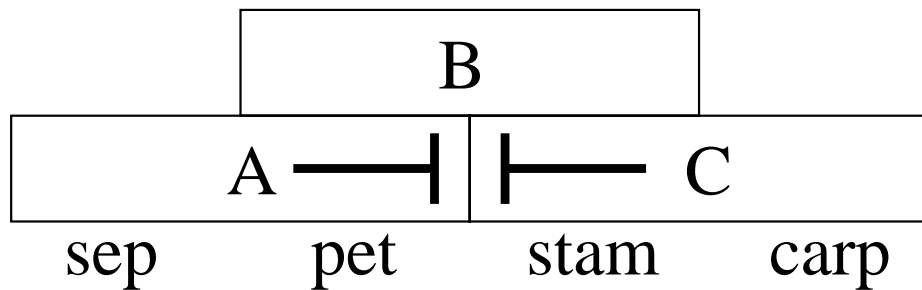
AP1 mRNA in  
ag mutant plants  
In all whorls



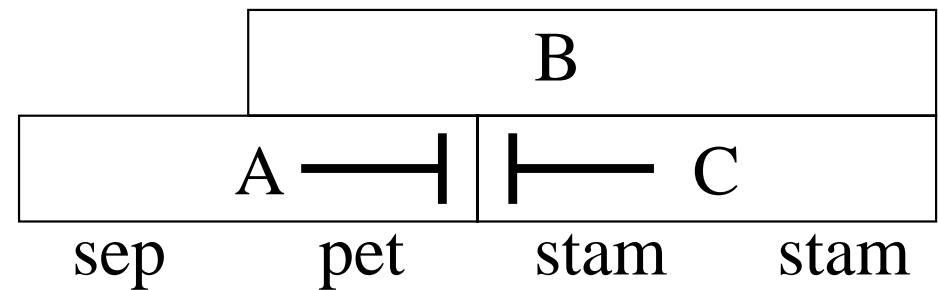
## Further additions to the ABC model : SUPERMAN



Wild type

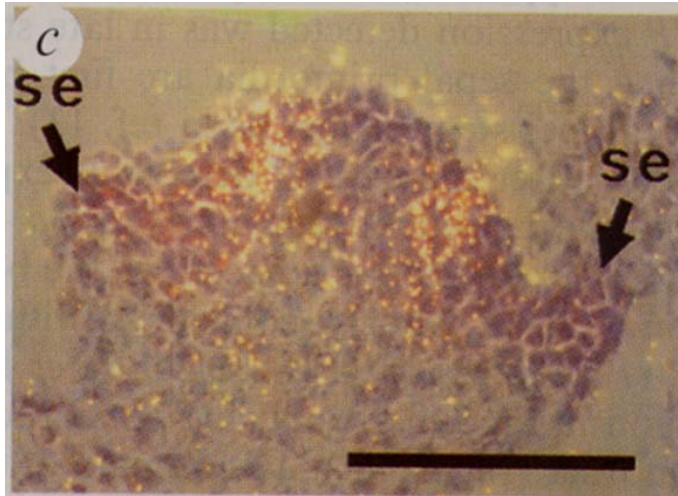


*superman*

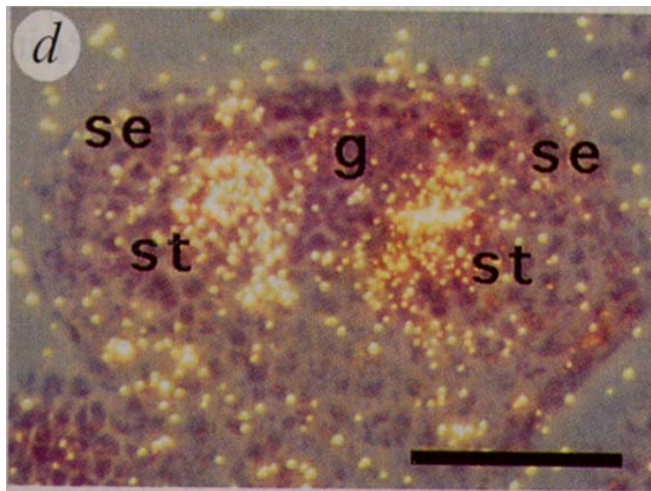


## **SUPERMAN is expressed at the boundary between the third and fourth whorl**

Stage 3



Stage 4

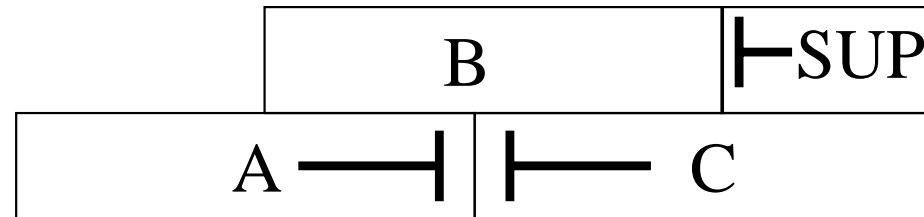


SUP encodes a likely zinc-finger DNA-binding protein containing Cys2 His2 type zinc fingers and a Serine/proline rich activation domain.

# ABC model

## Summary

Wild type

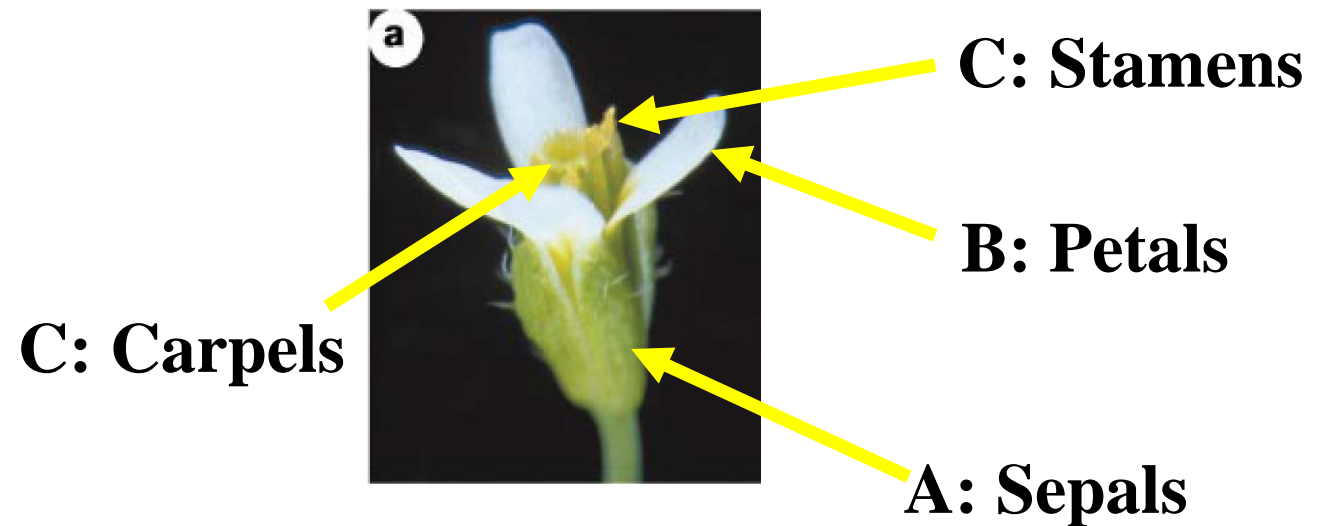


sep

pet

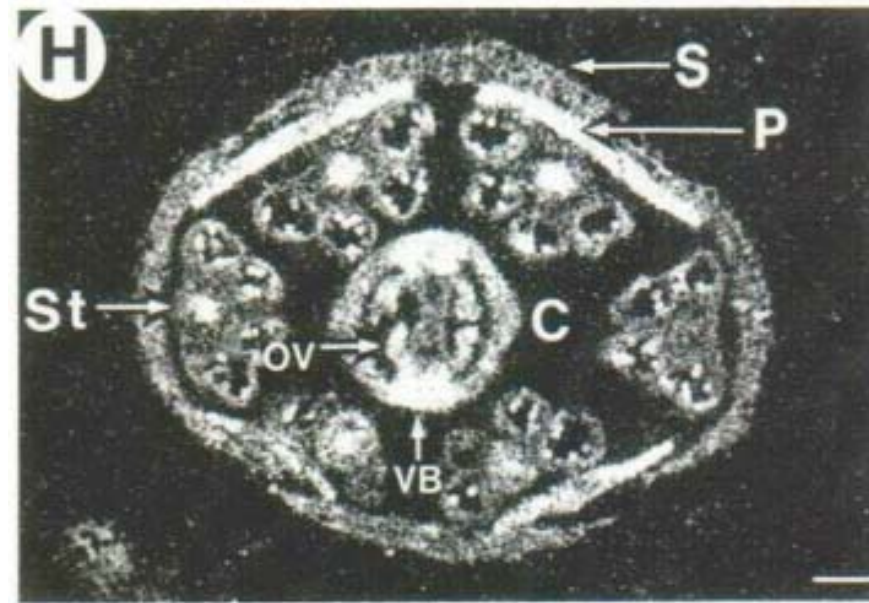
stam

carp

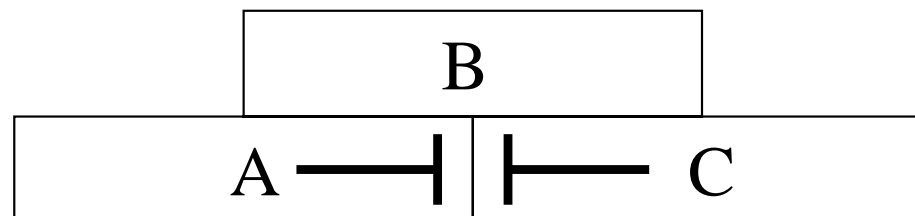


**APETALA2 is a class A gene required for whorl 1 and 2,  
but is expressed in all whorls.**

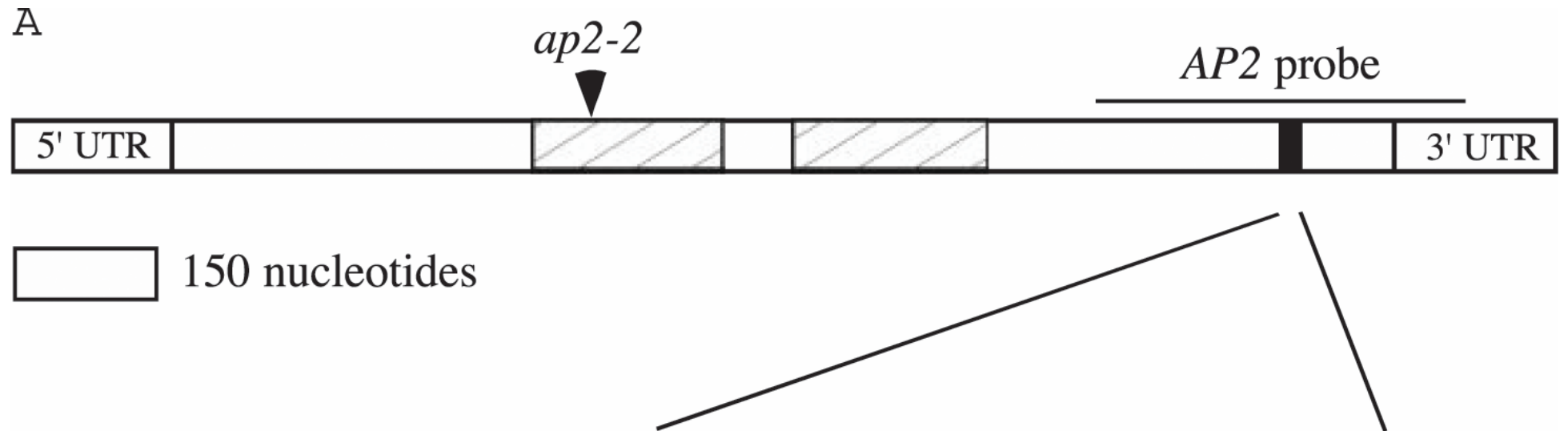
**How is AP2 activity restricted to whorls 1 and 2 ?**



Wild type



# Arabidopsis microRNA172 has homology to the AP2 gene



wild-type <i>AP2</i> RNA	5' gcu gca gca uca uca gga uuc ucu 3'
miRNA172a-1, -2	3' ua <b>cgu</b> <b>cgu</b> <b>agu</b> <b>agu</b> <b>ucu</b> <b>aag</b> a 5'
miRNA172b-1, -2	3' <b>ga</b> <b>cgu</b> <b>cgu</b> <b>agu</b> <b>agu</b> <b>ucu</b> <b>aag</b> a 5'
miRNA172c	3' ua <b>cgu</b> <b>cgu</b> <b>agu</b> <b>agu</b> <b>ucu</b> <b>aag</b> g 5'



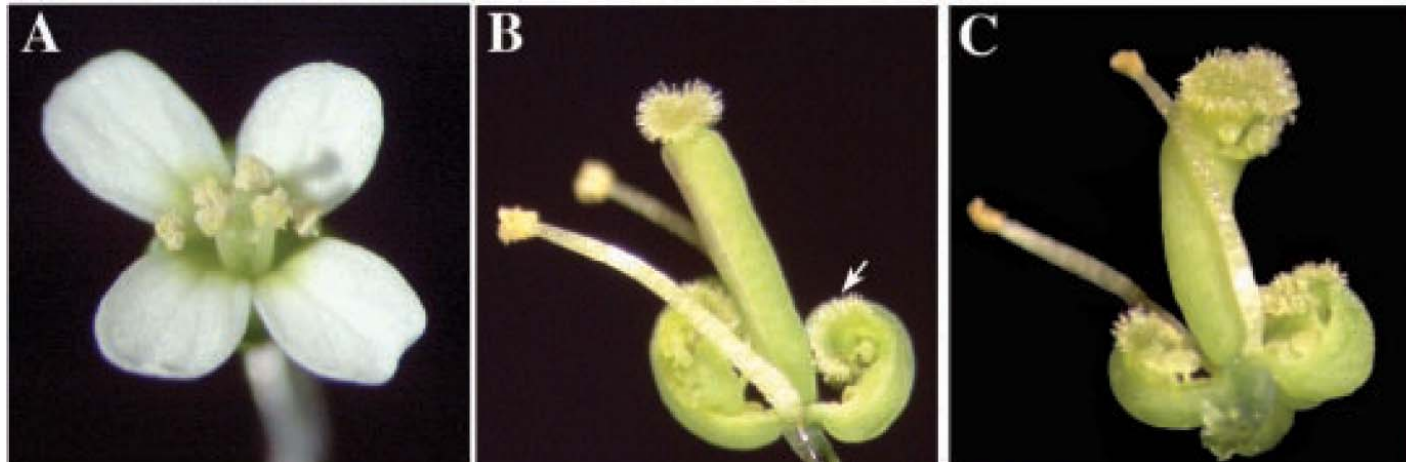


**Expression of mutant forms of MIR172 from  
a viral promoter  
causes an *ap2* mutant phenotype**

**WT**

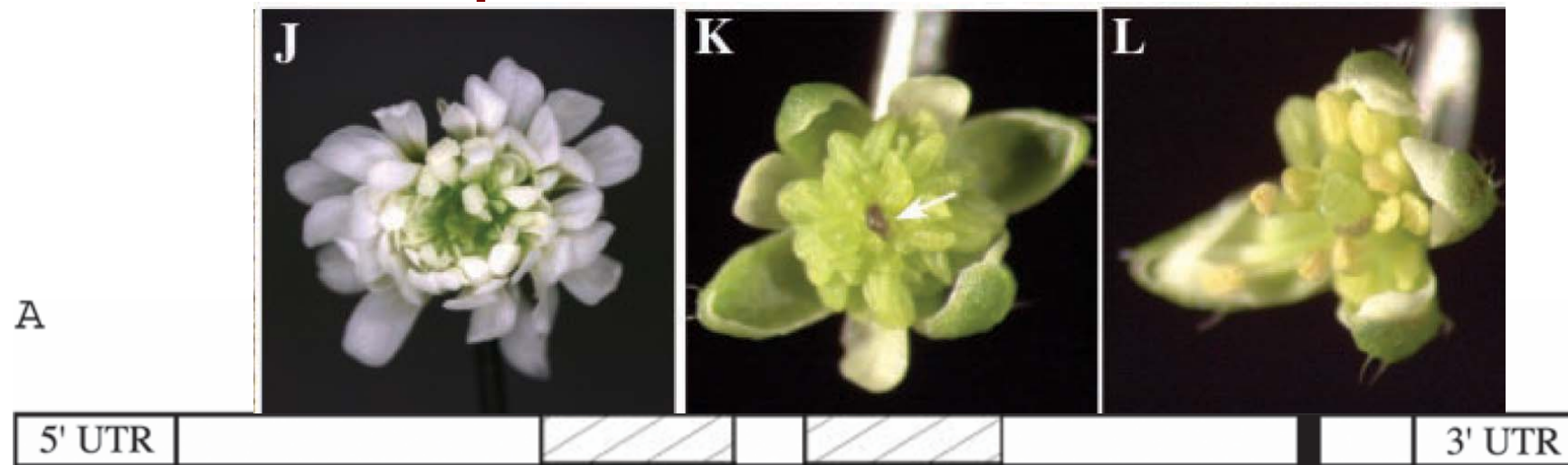
***ap2* mutant**

**35S::MIR172**



**Expression of MIR172 at high levels in all cells  
prevents AP2 function**

# Expression of a mutant form of AP2 mRNA that has reduced homology to MIR172 causes increased petal number and more floral whorls



150 nucleotides

AP2 protein

*AP2m3* RNA

*AP2m1* RNA

wild-type *AP2* RNA

miRNA172a-1, -2

miRNA172b-1, -2

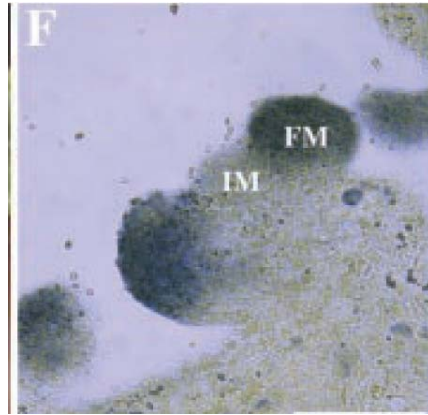
miRNA172c

	A	A	A	S	S	G	F	S	
5'	gca	gcu	gcu	ucc	ucu	ggu	uuc	ucu	3'
5'	gca	gcu	gcu	ucc	uca	ggu	uua	ucu	3'
5'	gcu	gca	gca	uca	uca	gga	uuc	ucu	3'
3'	ua	cgu	cgu	agu	agu	ucu	aag	a	5'
3'	ga	cgu	cgu	agu	agu	ucu	aag	a	5'
3'	ua	cgu	cgu	agu	agu	ucu	aag	g	5'

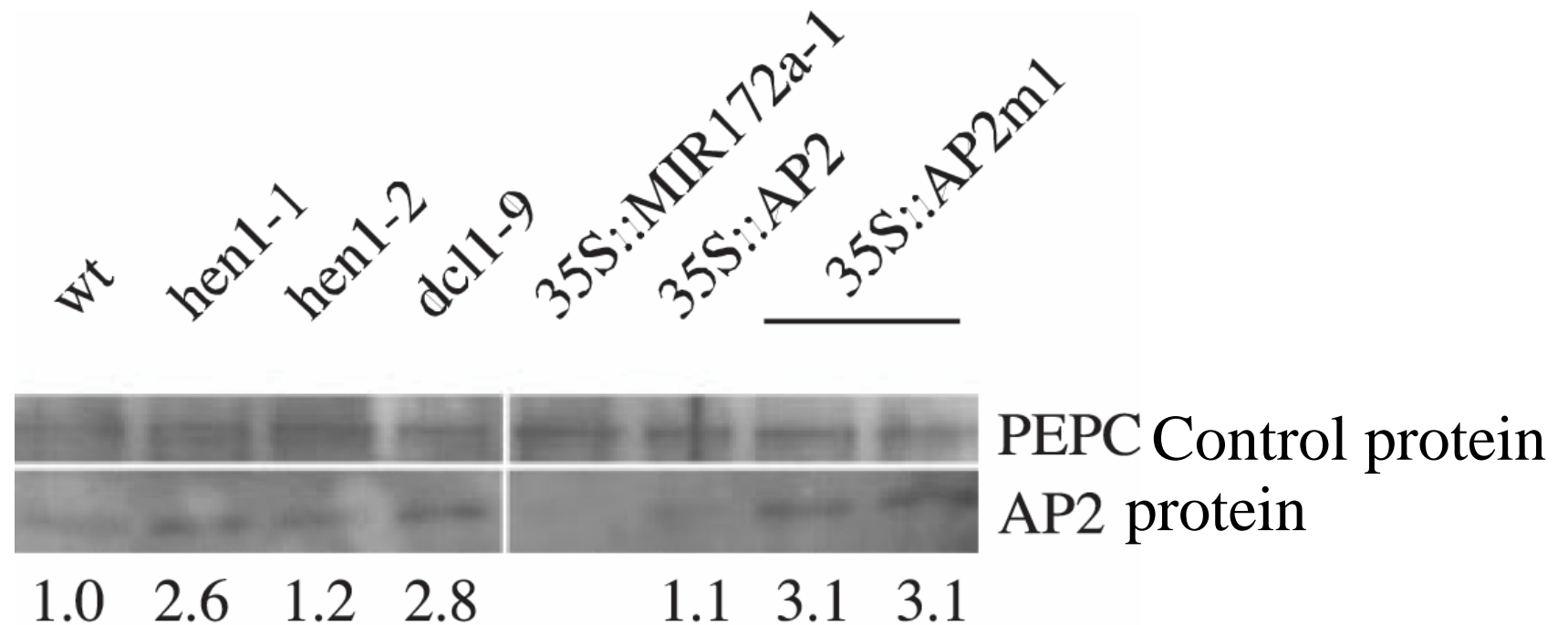


# MIR172 is expressed only in the inner whorls in older floral primordia and reduces AP2 protein levels

Floral meristems

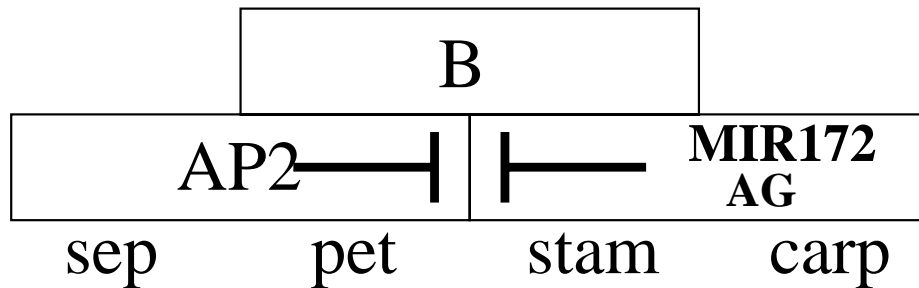


Older, Stage 7 flower

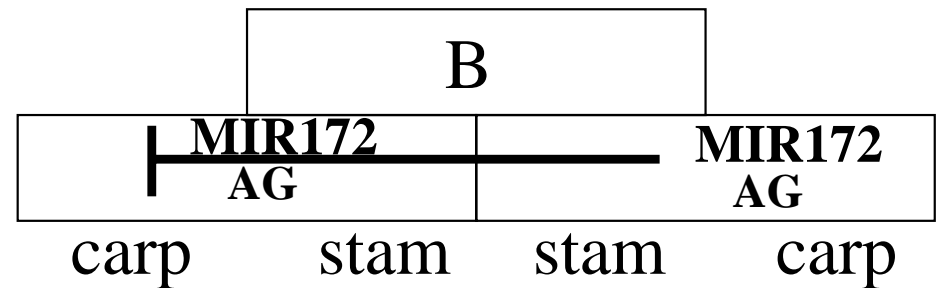


**MIR172 restricts AP2 activity to the 1 and 2 whorl  
so that AG expression is prevented in these whorls  
but can occur in whorls 3 and 4**

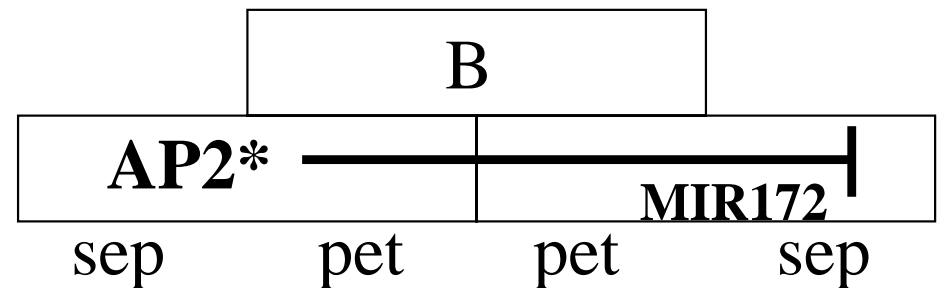
**Wild type**



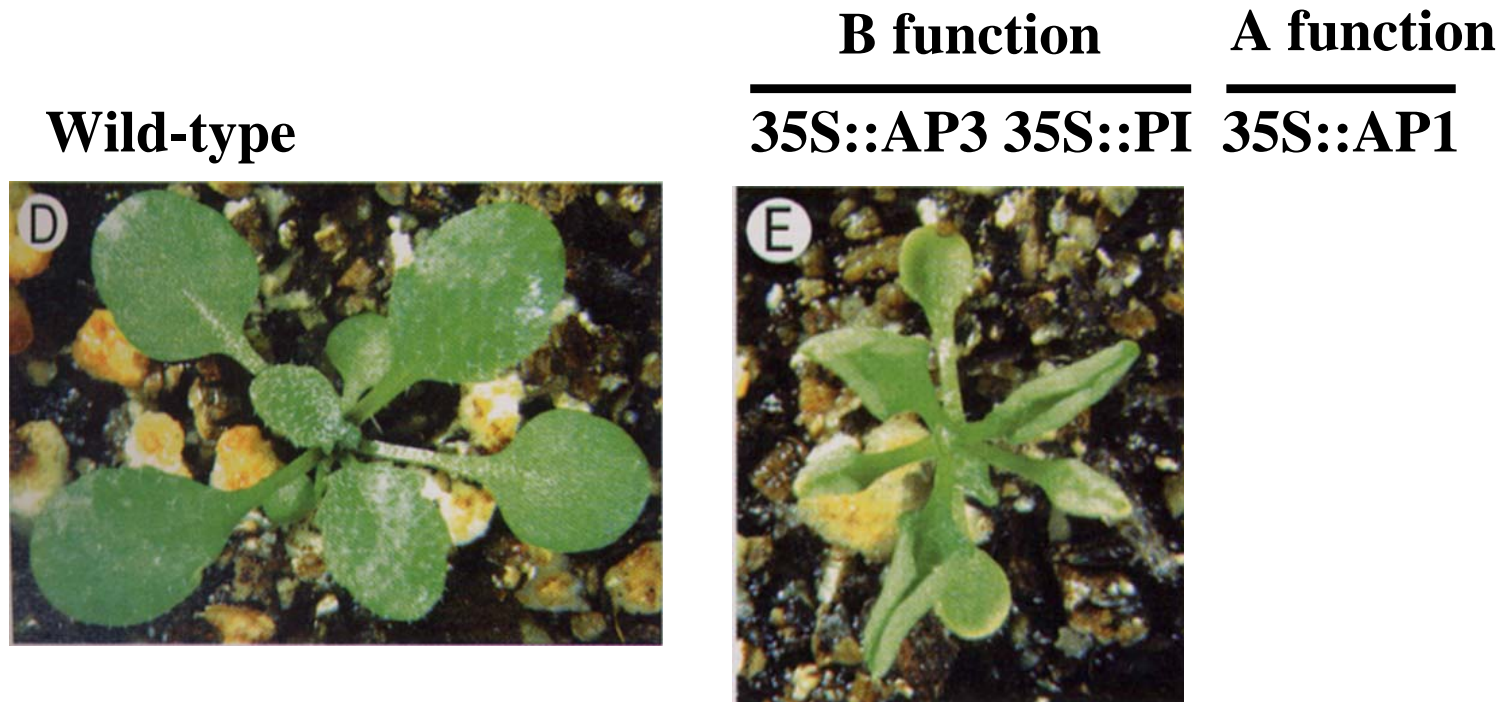
**35S::MIR172**



**35S::AP2\* not recognised by MIR172**



# Misexpression of B or A and B in leaves is not sufficient to convert leaves to petals



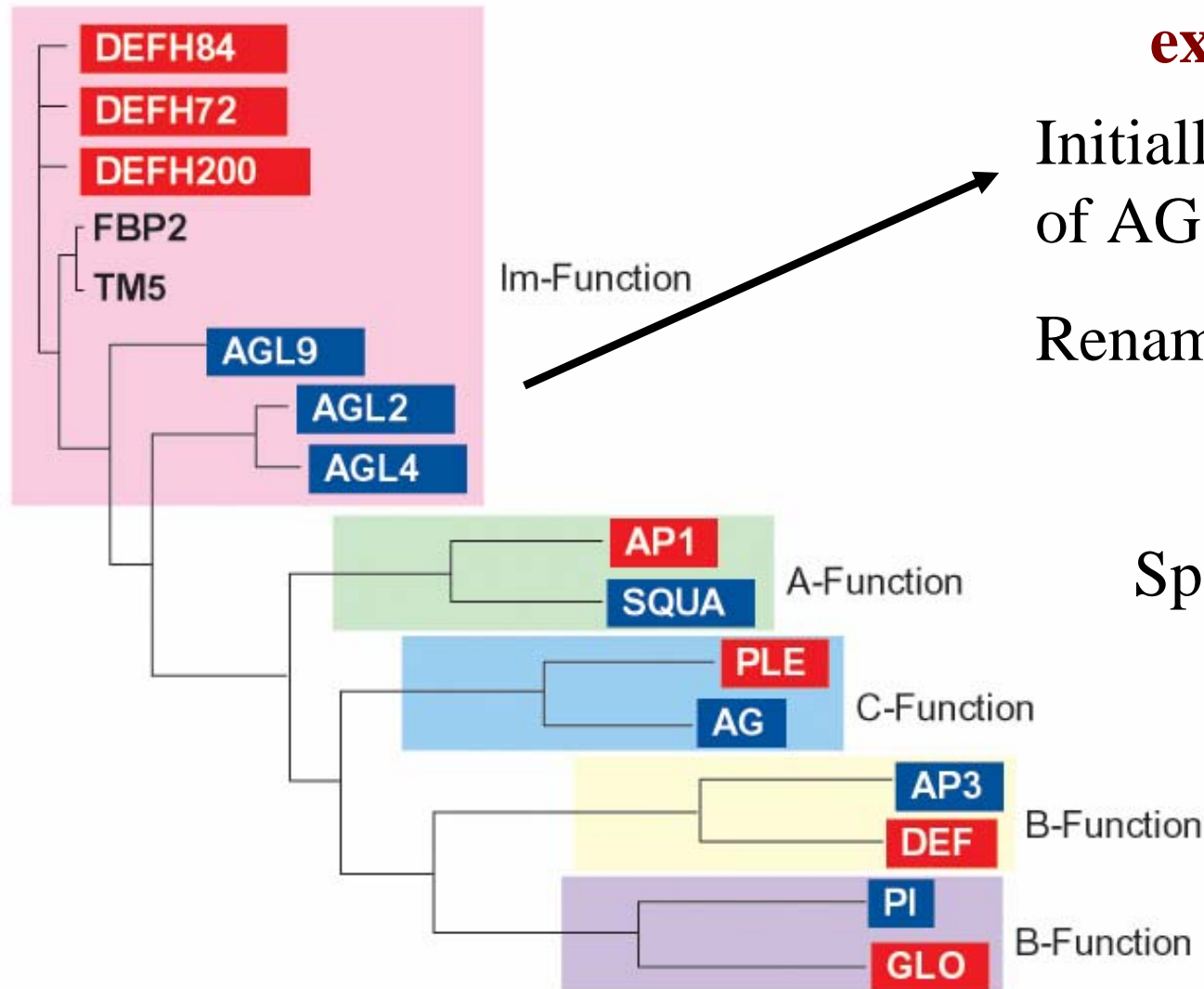
Suggests another floral-specific factor is absent in the leaves.

This was not identified by initial genetic screens

## More MADS box genes expressed in the flower

Initially identified as homologues  
of AG – AGL2, AGL4, AGL9

Renamed sepallata 1,2,3



Specific expression patterns –  
in whorls 2,3,4, although  
AGL2 and AGL9 are also  
expressed in whorl 1 of  
younger flowers.

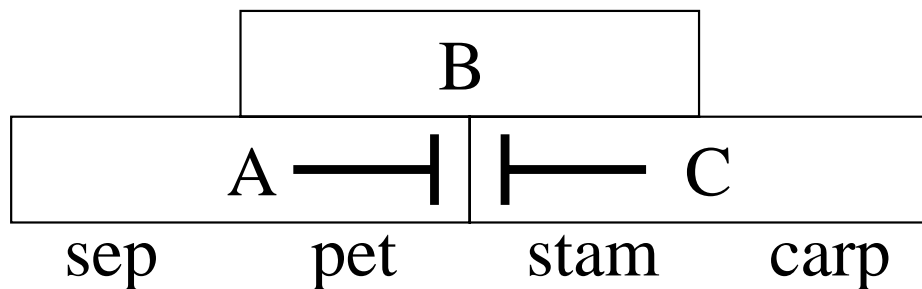
**Genes in red from *Antirrhinum***  
**Genes in blue from *Arabidopsis***

# Inactivation of SEP1 SEP2 and SEP3 in triple mutants

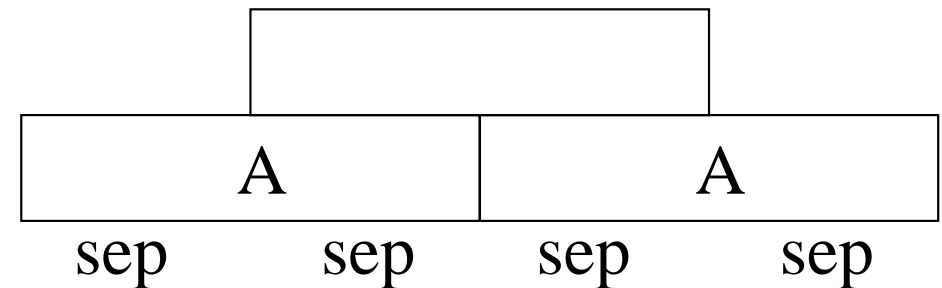


Were not identified in  
Original mutant screens  
Because of redundancy  
Between proteins.

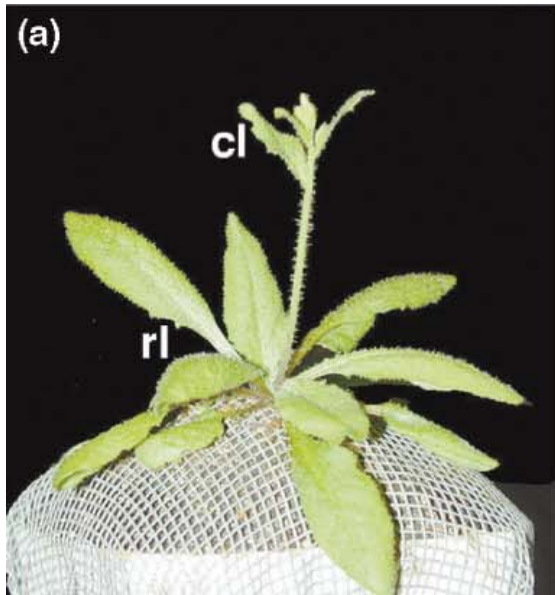
Wild type



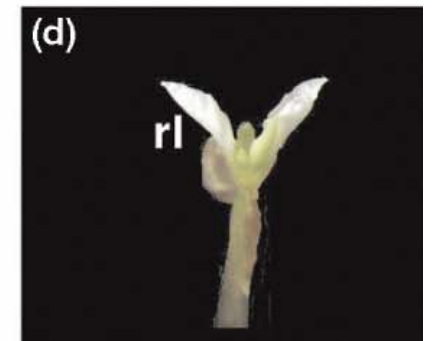
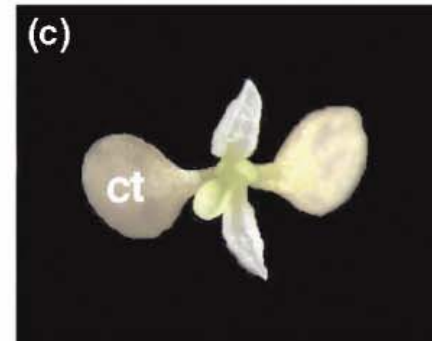
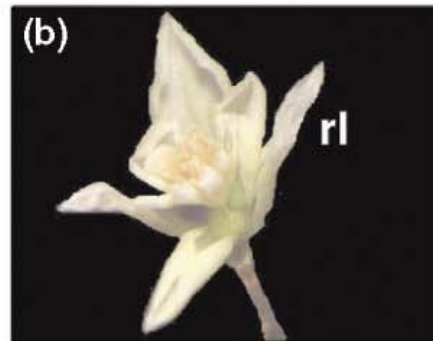
sep1 sep2 sep3



## Misexpression of SEP1 AP1 PI and AP3 in leaves creates petals

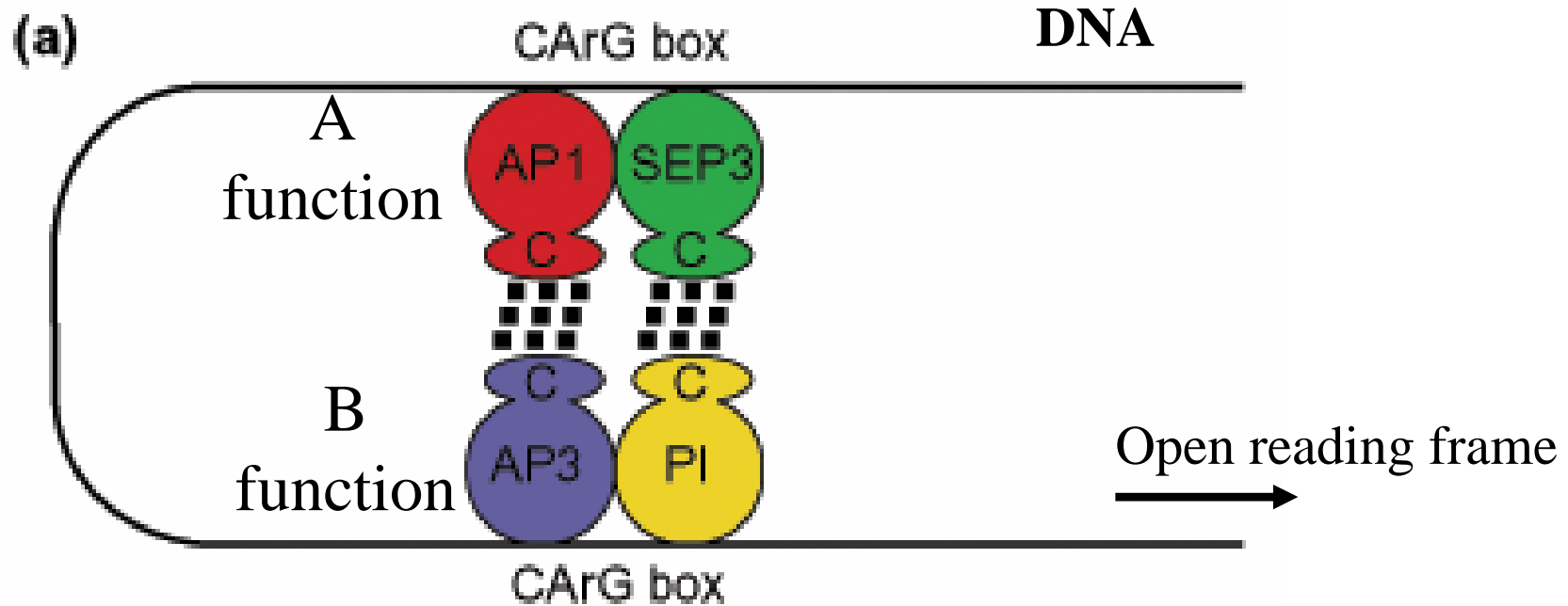


### WHOLE PLANTS

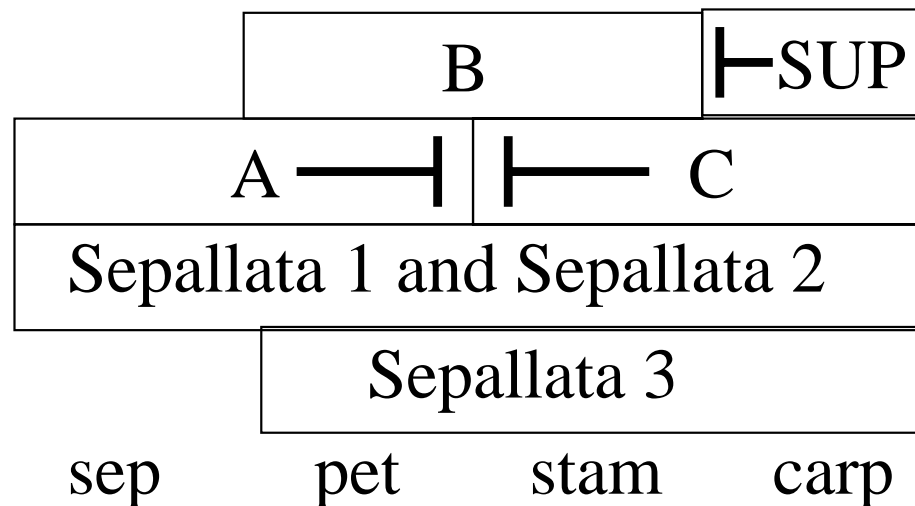


Plants carrying combination of  
35S::AP1 35S::SEP2 35S::PI 35S::AP3

**Therefore a combination of AP1 PI AP3 SEP is sufficient  
To confer B function.**

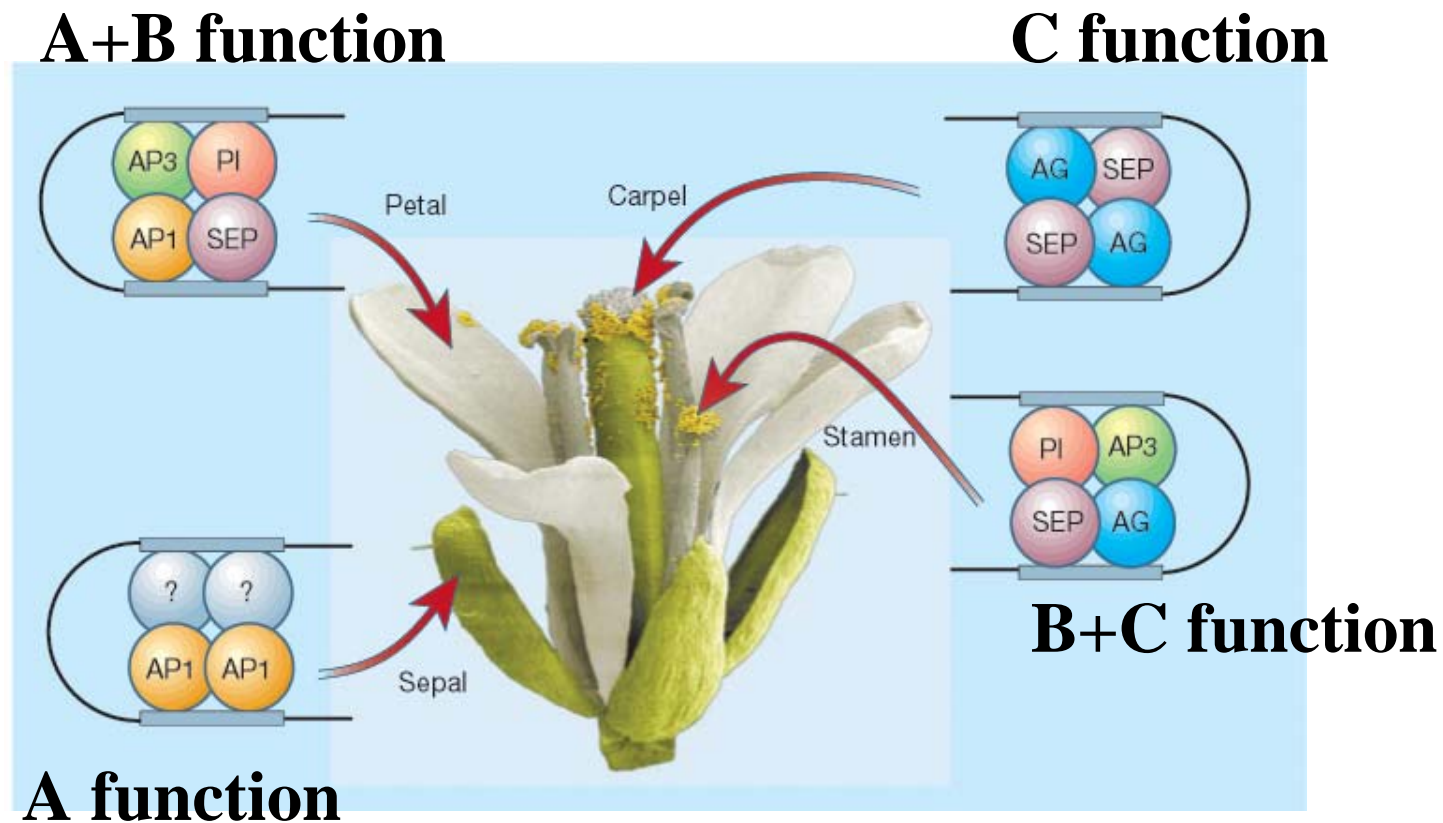


**Model explains how A function (AP1) and B function (AP3/PI) Combine to specify the second whorl – petals.**





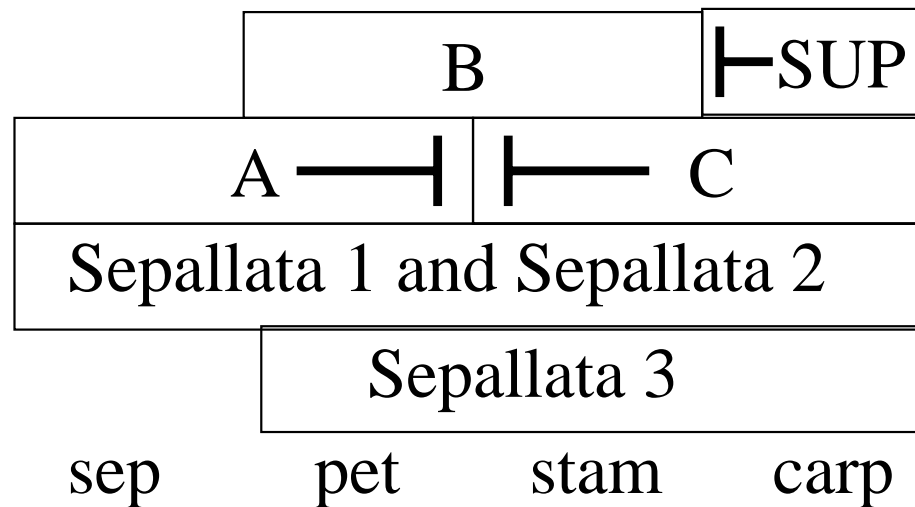
**Similar multimeric complexes of MADS box proteins are proposed to specify the other whorls**





# SEP proteins mediate multimeric complexes between PI/AP3 and AG or between PI/AP3 and AP1

Wild type



Formation of multimeric complexes suggests mechanism for combining  
A/B and B/C functions within the ABC model.

- this presentation

- PDFs

WEB ADDRESS:

[www.mpiz-koeln.mpg.de](http://www.mpiz-koeln.mpg.de)

Developmental Cell, Vol. 2, 135-142, February, 2002, Copyright ©2002 by Cell Press

## Building Beauty: The Genetic Control of Floral Patterning

Review

Jan U. Lohmann<sup>1</sup> and Detlef Weigel<sup>1,2,3</sup>

<sup>1</sup>Plant Biology Laboratory  
The Salk Institute for Biological Studies  
La Jolla, California 92037

<sup>2</sup>Department of Molecular Biology  
Max Planck Institute for Developmental Biology  
72076 Tübingen  
Germany

Floral organ identity is controlled by combinatorial action of homeotic genes expressed in different territories within the emerging flower. This review discusses recent progress in our understanding of floral homeotic genes, with an emphasis on how their region-specific expression is regulated.

about the mechanisms underlying this process. Because at this point there is a very large number of original publications in this field, we have cited reviews for most of the work published before the mid-1990s.

### The ABCs of Flower Development

Contemporary work on floral patterning began with the study of a series of mutants in which floral organs develop normally, but in the inappropriate whorl. Such mutants had been collected from garden snapdragon, *Antirrhinum majus*, by Hans Stubbe, and from the mustard relative *Arabidopsis thaliana* by Maarten Koornneef. In the late 1980s, three groups, headed by Enrico Coen in the United Kingdom, Elliot Meyerowitz in the United States, and Heinz Saedler in Germany, recognized the value of these mutants as homeotic mutants, and used them to initiate molecular and genetic studies of floral

Forschung

Abt. Entwicklungsbiologie de  
Pflanzen

George Coupland