

# B.2.16 is a non-lethal modifier of the $Dark^{82}$ mosaic eye phenotype in $Drosophila\ melanogaster$

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#### **Abstract**

Genetic screens have been used to identify genes involved in the regulation of different biological processes. We identified growth mutants in a Flp/FRT screen using the *Drosophila melanogaster* eye to identify conditional regulators of cell growth and cell division. One mutant identified from this screen, B.2.16, was mapped and characterized by researchers in undergraduate genetics labs as part of the Fly-CURE. We find that B.2.16 is a non-lethal genetic modifier of the  $Dark^{82}$  mosaic eye phenotype.

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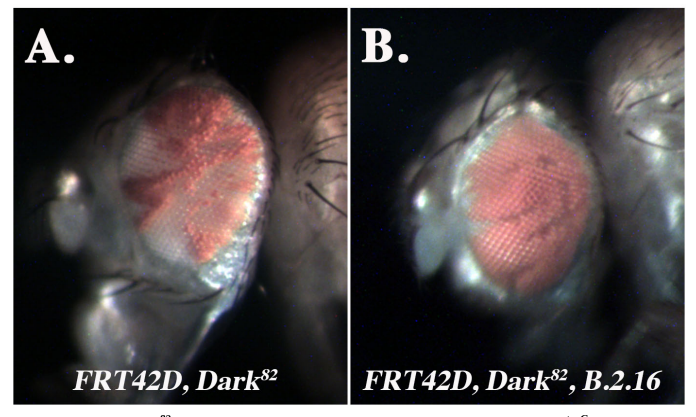


Figure 1. The B.2.16,  $Dark^{82}$  mosaic eye results in an increase in the ratio of mutant pigmented ( $w^{+mC}$ ) tissue compared to wild type unpigmented tissue: A.  $Dark^{82}$  mosaic eye (control), mutant tissue is pigmented. B.  $Dark^{82}$ , B.2.16 mosaic eye, mutant tissue is pigmented.

# **Description**

Table 1: Complementation tests conducted with mutant B.2.16

2R deficiency kit overlapping <i>Dark</i>					
Deficiencies tested within Dark deficiency region					
Deficiency	BDSC Stock #	Region	Complementation test with <i>B.2.16</i>		
Df(2R)BSC382	24406	2R: 16,776,16416,901,625	Complement		
Df(2R)CG15614 <sup>attP</sup>	84470	2R: 16,945,10716,947,153	Complement		
Df(2R)ED1	6916	2R: 17,026,72717,097,322	Complement		
Df(2R)BSC433	24937	2R: 17,062,91517,097,315	Complement		
Single genes tested within	Single genes tested within Dark deficiency region				
Gene	BDSC Stock #	Allele	Complementation test with <i>B.2.16</i>		
Ehbp1	10931	Ehbp1 <sup>k09837</sup>	Complement		
CG8963	12432	CG8963 <sup>BG00665</sup>	Complement		
Pkc53E	20790	Pkc53E <sup>EY14093</sup>	Complement		
mute	22655	mute <sup>EY22147</sup>	Complement		
PIG-V	32781	PIG-V <sup>MI01787</sup>	Complement		
AsnRS-m	10671	AsnRS-m <sup>k07408</sup>	Complement		

CG6805	78974	CG6805 <sup>CR00657</sup> -TG4.0	Complement	
CG30460	59338	CG30460 <sup>MI134331</sup>	Complement	
Genes tested outside of 2R deficiency kit				
Gene	BDSC Stock #	Allele	Complementation test with B.2.16	
<b>Gene</b> <i>l(2)41ab</i>		<b>Allele</b> <i>l</i> (2)41Ab <sup>KV00249</sup>	Complementation test with <i>B.2.16</i> Complement	

In order to identify genes that contribute to the genetic regulation of *Drosophila* eve development and the regulation of tissue growth, an ethyl methanesulfonate (EMS) mutagenesis screen was performed utilizing the FLP/FRT system, which generates genetically mosaic clones in a tissue specific manner. The screen was performed starting with a copy of chromosome 2R carrying an allele of Death-associated APAF1-Related Killer (Dark), Dark<sup>82</sup>, which was generated by imprecise P-element excision and therefore retains eye pigmentation due to the presence of an insertion encoding mini-white (Akdemir et al. 2006). When homozygous, the Dark82 allele blocks cell death and allows for the detection of conditional regulators of eye development and tissue growth that might otherwise induce apoptosis (Kagev et al. 2012). Flies that bear a FRT42D, Dark<sup>82</sup> chromosome (2R) were mutagenized with EMS and then crossed to ey-Flp; FRT42D flies. The mosaic eyes of resulting offspring were analyzed for phenotypes associated with overall head and eye size, ratio of mutant to wild-type tissue (red over white), or developmental patterning defects (Kagey et al. 2012). One mutant from this screen, B.2.16, was selected for study here. The control mosaic eye, *Dark*<sup>82</sup>, has an average ratio of 50-60% mutant (pigmented) tissue (**Figure 1A**). In comparison the B.2.16,  $Dark^{82}$  mosaic eye ranges from 75-80% mutant tissue ( $w^{+mC}$ ) demonstrating a consistent increase in the amount of mutant tissue from the *Dark*<sup>82</sup> mosaic ratio (**Figure 1B**). To understand the mechanism driving this increase in mutant tissue overrepresentation, *B.2.16* was mapped via complementation mapping by undergraduate researchers who were part of the Fly-CURE consortium. This type of mapping has been successful for previous mutants from this screen such as *Egfr*, *Ptc*, and *Shn* (Bieser et al. 2019, Kagev et al. 2012, Stamm et al. 2019)

The genetic mapping of B.2.16 on chromosome 2R was completed by four independent groups of undergraduate researchers at Illinois State University, Nevada State College, the University of Evansville, and the University of Detroit Mercy as a part of the Fly-CURE consortium (Bieser et al. 2018, Bieser et al. 2019, Stamm et al. 2019). Complementation tests were conducted using the 87 deficiency stocks from the Bloomington Stock Center 2R Deficiency Kit that are distal to the FRT42D location (Cook et al. 2012). With the hypothesis that the B.2.16 mutation is homozygous lethal, we mapped homozygous lethality by performing complementation tests using the 87 deficiency stocks. Virgin FRT42D, B.2.16, Dark82/CyO females were crossed to males from each of these deficiency stocks and the F1 progeny were scored for complementation. None of the deficiency stocks failed to complement the B.2.16 mutation. This finding suggests three possible hypotheses: 1) The B.2.16 mutation is linked with the *Dark*<sup>82</sup> mutation and therefore unable to be mapped via complementation due to the homozygous lethal nature of the *Dark*<sup>82</sup> allele. Two deficiencies from the deficiency kit (*Df*(2*R*)*ED2747* and *Df*(2*R*)*BSC*331) fail to complement *Dark*<sup>82</sup>; if the *B.2.16* mutation lies within this region of 2R:16,869,330..17,097,303, then a failure to complement for the *Dark*<sup>82</sup> allele would be indistinguishable from a failure to complement for the B.2.16 allele (Gramates et al. 2017). 2) The B.2.16 mutation lies in a region that is not covered by a deficiency stock. There are 46 predicted genes on chromosome 2R that are not covered by the deficiency kit (https://bdsc.indiana.edu/stocks/df/dfkit-info.html, Cook et al. 2012). If the mutation exists in one of these genes, it cannot be mapped using the deficiency kit. 3) Lastly, the *B.2.16* mutation is not a homozygous lethal mutation and therefore cannot be mapped via complementation tests.

We conducted additional experiments to investigate these hypotheses. To test the first hypothesis that the B.2.16 mutation is tightly linked to the lethal  $Dark^{82}$ , we set up complementation tests with smaller deficiency stocks and all available individual gene lethal alleles within the  $Dark^{82}$  region (see Table 1 for a list of stocks tested). We found that B.2.16 complemented all of these crosses within the  $Dark^{82}$  region, suggesting that B.2.16 is not a lethal mutation tightly linked to Dark. To test the second hypothesis that B.2.16 lies in a region of 2R that is not covered by the deficiency kit, we investigated the 46 predicted genes that are not covered by the Df kit. Of these predicted genes, 33 are considered either pseudo-genes or 'gene model not supported' on FlyBase and therefore are unlikely candidates (Thurmond  $et\ al.\ 2019$ ). For the remaining 13 genes, there were two available lethal alleles, which were tested and shown to complement with B.2.16 (Table 1). Since our testing for complementation is limited to available lethal alleles and deficiencies, we also investigated genome-wide sequencing (GWS) data for B.2.16 to look for changes in any gene not directly tested via complementation. We found no mutations within these



untested genes (either the *Dark* overlap region or in areas not covered by the Df kit) that would truncate or alter protein function, per SNP effect predictions by SnpEff software (Cingolani *et al.* 2012).

Given that the B.2.16 mutation complemented all stocks tested across 2R (Df Kit, additional Df stocks, and single allele stocks) and that GWS provides no clear candidate mutations within the genes not tested by complementation, we conclude that the B.2.16 mutation is a non-lethal mutation that modifies the  $Dark^{82}$  mosaic eye phenotype resulting in an increased ratio of mutant tissue to wild-type tissue.

## Reagents

FRT42D Dark<sup>82</sup>/CyO (Akdemir et al.., 2006) FRT42D Dark<sup>82</sup>, B.2.16 /CyO (this manuscript)

Ey-Flp; FRT42D (BDSC 8211)

Bloomington Drosophila Stock Center 2R Deficiency Kit (Cook et al., 2012)

Additional Bloomington Stocks (See Table 1 for complete list of stock numbers)

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