RESEARCH ARTICLE

The Gene Encoding Dihydroflavonol 4-Reductase Is a Candidate for the *anthocyaninless* Locus of Rapid Cycling *Brassica rapa* (Fast Plants Type)

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Abstract

Rapid cycling *Brassica rapa*, also known as Wisconsin Fast Plants, are a widely used organism in both K-12 and college science education. They are an excellent system for genetics laboratory instruction because it is very easy to conduct genetic crosses with this organism, there are numerous seed stocks with variation in both Mendelian and quantitative traits, they have a short generation time, and there is a wealth of educational materials for instructors using them. Their main deficiency for genetics education is that none of the genetic variation in RCBr has yet been characterized at the molecular level. Here we present the first molecular characterization of a gene responsible for a trait in Fast Plants. The trait under study is purple/nonpurple variation due to the *anthocyaninless* (*ANL*) locus, which is one of the Mendelian traits most frequently used for genetics education with this organism. We present evidence that the DFR gene, which encodes dihydroflavonol 4-reductase, is the candidate gene for the *anthocyaninless* (*ANL*) locus in RCBr. DFR shows complete linkage with *ANL* in genetic crosses with a total of 948 informative chromosomes, and strains with the recessive nonpurple phenotype have a transposon-related insertion in the *DFR* which is predicted to disrupt gene function.

Introduction

Rapid cycling *Brassica rapa* (RCBr), more commonly known as Wisconsin Fast Plants, is a widely used organism in science education. They are used at both K-12 and college levels as a model system for topics including growth and development, ecology, physiology, reproduction, and genetics. They have a six week generation time, a petite growth habit, easily grown in conventional potting soil, and do not require greenhouses or growth chambers. They thrive at room temperature under ordinary fluorescent lights, although the light must be constant and intense [1]. Their other major strength is that a huge volume of educational modules and support material for teachers has been developed by the Wisconsin Fast Plants Program [2].

Fast Plants are an appealing system for teaching genetics. Because *Brassica rapa* are self-incompatible for pollination, a student at any level can conduct a genetic cross by simply using...
a cotton swab to transfer pollen between plants [1]. A wide collection of seed stocks is available for experimentation in both Mendelian and quantitative traits, and these are readily available to educators from Carolina Biological Supply (Burlington, NC, USA) or from the Rapid Cycling Brassica Collection [2]. Dozens of Mendelian traits have been identified but the most frequently used are anthocyanin expression (purple versus non-purple stem), leaf color (green versus yellow-green), and trichome (hairless versus presence of hairs). Pigmentation and trichome density also exhibit quantitative variation [3]. Among plants that express anthocyanin pigments there is great variation in the intensity of the pigmentation and this variation is controlled by both polygenic inheritance and environment [4,5]. However, despite their widespread use in genetics education, none of the heritable traits in Fast Plants strains have been characterized at the molecular level.

The molecular genetics of anthocyanin expression has been studied in many plant systems. Anthocyanins are a group of flavonoid pigments that give color to flowers, fruit, stems, and leaves to varying degrees in many plant species. They play diverse roles including, but not limited to, attracting pollinators and frugivores, protection from herbivores, protection against ultraviolet light, and scavenging free radicals [6,7]. Numerous “anthocyanin genes” have been identified in plants, including the Brassicaceae family. Anthocyanin genes fall in to two broad categories: “structural genes” that encode enzymes in the biosynthetic pathways leading to formation of anthocyanins and “regulatory genes” that encode transcription factors needed for activation of the structural genes [8,9].

Here we present the first molecular characterization of a gene responsible for a trait in Fast Plants. We report here that non-purple strains of RCBr have a transposon-related insertion in the fourth exon of the gene encoding dihydroflavonol-4-reductase (DFR). This mutation shows complete linkage with the anthocyaninless (anl) locus indicating that anl and DFR are most likely the same gene.

Materials and Methods

Plant Strains

The rapid cycling Brassica rapa (Fast Plants type) strains Purple Stem, Hairy (ANL/ANL) and Non-Purple Stem, Yellow-Green Leaf (anl/anl) were obtained from Carolina Biological Supply (Burlington, NC). Purple (ANL/ANL) strain DWRCBr76 was reported previously [10]. Non-purple (anl/anl) strain DWRCBr57 was derived from DWRCBr53 [10] with selection for homozygosity of DNA markers on chromosome A09.

Genetic Marker Developments

Brassica rapa genome sequence was obtained from the BRAD database [11] or from Phytozome [12]. Short tandem repeat sequences (STR) were identified using Tandem Repeats Finder [13]. Primers for amplification of STR were designed using Primer-BLAST [14] and tested for polymorphism between DWRCBr76 and DWRCBr57. Genetic markers usable for mapping are given in Table 1.

DNA Purification

For DNA purification, plant tissue was disrupted in a ground glass homogenizer in 500 μl of 50 mM tris base, 240 mM NaCl, 25 mM EDTA, 0.5% SDS with RNAse H. The homogenate was extracted with chloroform and isoamyl alcohol (24:1). DNA was precipitated with an equal volume of isopropyl alcohol. The pellet was washed with 70% ethanol, air dried, and dissolved in TE buffer, pH 7.5.
Polymerase Chain Reaction (PCR)
Genomic DNA was amplified by PCR with the following protocol: 94° for 2 min., 30 cycles of 94° for 30 sec., 61° for 1 min., 72° for 1 min., and finally 72° for 4 min.

Genotyping
For most genetic markers, alleles were resolved on 1.2% agarose gels. Markers that could not be resolved on agarose gels were resolved in nondenaturing polyacrylamide gels consisting of 8% total acrylamide (19:1 acrylamide:bis) in Tris-Borate EDTA buffer.

Mapping by Backcross
Strains DWRCBr76 and DWRCBr57 were crossed and the F1 generation was backcrossed to DWRCBr57. BC1 seeds were sprouted on damp blotter paper under a bank of six 40W fluorescent tubes. After three to five days of constant light exposure, a total of 126 seedlings were scored for phenotype (purple or non-purple) relative to seedlings of the parental strains. Tissue was collected for DNA purification. Genetic map distances in Kosambi centimorgans were determined using MapDisto [15]. Marker order was set by their position in the Brassica rapa genome FPsc ver. 1.3 in Phytozome [12].

Homozygosity Mapping
Strains DWRCBr57 and DWRCBr76 were crossed and pairs of F1 generation plants were intercrossed to produce an F2 generation. F2 seeds were sprouted as described above and only non-purple seedlings were collected for mapping. A total of 1199 F2 seedlings were produced. Of these, 402 were non-purple and used for homozygosity mapping. All 402 non-purple F2 seedlings (sprouted as described above) were genotyped for markers D9BrapaS7 and D9BrapaS6.

| Marker      | A09 Position1 | Forward Primer | Reverse Primer             |
|-------------|---------------|----------------|-----------------------------|
| D9BrapaS12  | 7,222,274     | CCAGCCAAATCGTCACTCATGCGA | TGCACTGCCTAAAGAATTGGAATACAC |
| D9BrapaS8   | 10,306,916    | AACAGGAGCTCCTCCATGCGG   | ACTCTCTATCGTTGGGCAAA   |
| D9BrapaS7   | 10,989,431    | CTCCTCTGCTGATTGTTG      | AATCCCGGCTAGTGGAGAAC |
| D9BrapaS9   | 11,459,243    | ACAGGGGAAGGAGCAGACTTT   | GAGATGACTCGCCCGGAACCC |
| D9BrapaS10  | 11,617,011    | GATCGGGCACTAGCCAAAAAA   | CACTTTACTAGGAAATGGTCACA |
| D9BrapaS11  | 11,877,226    | AATTCGAGCTAGGTCTGCTG    | AAGAAAGGCTTCCGCGCAATA |
| D9BrapaS15  | 11,890,565    | AGAGAATGAGAGGACAGGTGGA  | CCTCAACCTCTGACTTGACCT |
| D9BrapaS14  | 12,102,596    | GAGAGAGCAGGATGACG       | AGAGGAAAGCATGAGAGAGG |
| D9BrapaS13  | 14,111,236    | CCAAGAGGCAAGACTATTTAAA  | TTACAGCAACACACCAAGATA |
| D9BrapaS12  | 19,689,771    | AAGGAGGCTAGCTAGGTAAGAACA| AAGGCAATGCTAAGTCTTGCC |
| D9BrapaS6   | 19,474,795    | ATGGGCGGCTCTGTATCCA     | CTCCGGAGCTACCTGTCTGC |
| D9BrapaS5   | 29,261,082    | AGGTAAAGGCACTAACTGAAA   | TGGATATGTTGCTGCCCTGA |
| Bn9A3       | 30,357,828    | GAGGATCATCCCGAAGAAGGAAAG| CTTGGGAACATGAGAGATAG |
| D9BrapaS42  | 35,598,603    | TGGAGGCGTCAGGAGGACTGCTGA| AGCGATGAGACACCGAGTCCA |
| Park9HaeII2 | 36,389,743    | TGGCGGAGAAGAACACAGC     | TCCTAGCAGCTTCTACACCTC |

1Position of 5’ end of forward primer in Brassica rapa FPsc v1.3, DOE-JGI, http://phytozome.jgi.doe.gov/ [12].
2Source Slankster et al [10]
3Developed by Kresovich and applied to RCBr as reported by Slankster et al [10]
D9BrapaS5 and D9BrapaS12 are dominant markers which give a PCR product in DWRCBr76 but not in DWRCBr57.

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Individuals that were recombinant between these two markers were genotyped for additional markers within the interval.

**Molecular Cloning**

PCR amplicons were purified for DNA sequencing using a MinElute PCR Cleanup Kit (Qiagen Inc., Valencia, CA). Fragments were cloned into pGEM-T vector (Promega, Madison, WI, USA) and transformed into competent 10-beta *E. coli* cells (New England Biolabs, Ipswich, MA, USA).

**DNA sequencing**

Sequencing reactions were performed using ABI BigDye Terminator v3.1 Cycle Sequencing Kit and analyzed using the Applied Biosystems ABI Prism 3730 DNA Analyzer at the Wayne State University Applied Genomics Technology Center (agtc.wayne.edu). Nucleotide sequences were submitted to Genbank under accessions numbers KX185527, KX347549, and KX379243.

**Results**

**Fine mapping of *anthocyaninless* to proximal chromosome A09**

The *anl* locus was previously mapped to *Brassica rapa* chromosome A09 [16], but with low resolution. To more precisely map the position of *anl* we developed additional markers (Table 1) and generated larger sample sizes. In a backcross with 126 progeny, we mapped *anl* to within the interval between markers *D9Brapa6* and *D9Brapa7* (Fig 1). This interval is 3.8 centimorgans on our genetic map and 8.4 Mb in the *Brassica rapa* genome assembly FPsc v1.3.

We next used homozygosity mapping to further refine the position of the *anl* gene. We genotyped 402 non-purple (*anl/anl* genotype) F2 generation seedlings (a total of 804 informative chromosomes) for markers *D9BrapaS6* and *D9BrapaS7*. Among these, 57 individuals were recombinant between one of the markers and the *anl* locus. Genotyping of these 57 recombinants for additional markers between *D9BrapaS6* and *D9BrapaS7* (Fig 2) allowed us to narrow the position of *anl* down to a 212 kilobase interval which contained 28 predicted genes (Table 2). Of the annotated genes in the interval, Phytozome gene *Brara.I01754*, which encodes dihydroflavonol 4-reductase (*DFR*) stood out as the most likely candidate because mutations of this gene have been shown to result in an anthocyaninless phenotype in at least ten other plant species across the plant kingdom [9,17–25].

**Mutation in the *DFR* gene**

Analysis of the *DFR* gene in purple and nonpurple strains revealed that the nonpurple strains have an insertion mutation in the fourth exon of the *DFR* gene (Fig 3). PCR reactions with primers anchored in exon 4 (Table 3) of the gene indicated an insert of 300–400 bp (Fig 3-B).

We cloned genomic DNA of the *DFR* gene from wild type strains and mutant strains for nucleotide sequencing. Alignment of the nucleotide sequences of the *DFR* gene from wild type strains with the predicted mRNA of Phytozome gene *Brara.I01754* indicated that the purple strains have a functional *DFR* gene sequence. Initial attempts to sequence the mutant allele of the gene failed at a point roughly in the middle of predicted exon 4. This suggested a sequence that was forming a secondary structure such as a hairpin that is refractory to the sequencing reaction. Therefore we subcloned separate sections of the insert for sequencing. We screened a collection of restriction enzymes and found that one, *HpyCH4IV*, divided the segment approximately in half. We subcloned each half and were successful at sequencing each half separately. Assembly of these sequences revealed that the insertion sequence has a 340 bp sequence
Fig 1. Genetic linkage map of chromosome A09 including DNA markers tightly linked to the *anthocyaninless* (ANL) locus. Genotype data are from 126 BC1 progeny. Distances are in Kosambi centimorgans and all linkages have a LOD > 3.0.

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consisting of an inverted repeat with two 170 bp halves that form a perfect palindrome. The inverted repeat was flanked by direct terminal repeats (Fig 3-C). To verify that this sequence was not a cloning artifact, we designed a PCR primer that was anchored in the insert and designed to prime away from the center of the palindrome and used this to test for the existence of the palindrome in genomic DNA samples from the nonpurple strains. When the insert-specific primer was paired with a primer from the gene sequence upstream of the insert, it produced a PCR product. The insert-specific primer also produced a PCR product when paired with a primer from the gene sequence downstream of the insert (Fig 3-D). This confirmed the structure indicated by the DNA sequencing data.

The insertion introduces a premature stop codon into the gene. There is TGA beginning at the 53rd base of the palindrome which is in frame when aligned with the predicted mRNA of Brara I01754 (Fig 3-C). The presence of flanking direct repeats suggested a transposition event. A BLAST search of the Brassica rapa genome using the inserted sequence did not identify any other palindromic sequences, but it did identify over 100 sequences with greater than 90% identity and 90% length equivalent to one “half palindrome”. Of all of the BLAST hits in Brassica the longest were 170 base pairs. We did not find any other examples of the inverted repeat arrangement in either the FPsc v1.3 genome in Phytozome [12] or Chiifu-401 genome in BRAD [11]. A similar result was found in Brassica olearacea, but no such sequence was found in other species, including Arabidopsis thaliana. Thus the insertion in DFR found in non-purple rapid cycling Brassica rapa appears to be specific to its genus.

Complete linkage between anl and DFR
To confirm the identity of the DFR gene as the anthocyaninless locus, we genotyped the BC1 and F2 groups described above for the mutant and wild type alleles of DFR. Together these groups provide 930 informative chromosomes but there were no recombinants between DFR
and *anl*. We also genotyped 18 non-purple F2 progeny from a cross of Purple Hairy and Non-purple Stem, Yellow Green Leaf. These were also homozygous for the mutant allele of *DFR*.

Some purple plants have an insertion mutation in the last intron

Alignment of genomic DNA sequences of the *DFR* gene from the two purple strains Purple Stem, Hairy and DWRCBr76 revealed that the DWRCBr76 strain carries a 754 bp insertion in the intron between exons 4 and 5 of the *DFR* gene (Fig 4).

A simple test for the *anl* mutation suitable for instructional labs

With the PCR described for Fig 4-C one combination of primers specifically detects the mutant allele and another detects the wild type allele. Normally, to detect an insertion or deletion by PCR one simply uses a single pair of primers and observes a change in the size of the PCR product, and a heterozygote is detected as a double band. However, when we perform such a PCR with an ANL/anl heterozygote the wild type allele amplifies preferentially (Fig 5-A), presumably due to the palindromic insert reducing PCR efficiency. Therefore, each allele must be detected in separate PCR reactions. This strategy makes a very robust and clear PCR assay for

| Gene         | Annotation                                      |
|--------------|-------------------------------------------------|
| Brara.I01732 | auxin-inducible                                 |
| Brara.I01733 | none                                            |
| Brara.I01734 | solute carrier family 35                        |
| Brara.I01735 | 60S ribosomal protein L3-related                |
| Brara.I01736 | zinc finger FYVE domain containing protein      |
| Brara.I01737 | clathrin coat assembly protein                   |
| Brara.I01738 | uncharacterized protein family                   |
| Brara.I01739 | Agenet                                           |
| Brara.I01740 | Protein of unknown function, DUF617             |
| Brara.I01741 | Protein of unknown function, DUF547             |
| Brara.I01742 | Apoptosis-promoting RNA-binding protein          |
| Brara.I01743 | none                                            |
| Brara.I01744 | none                                            |
| Brara.I01745 | histone H3                                       |
| Brara.I01746 | uncharacterized protein ath:AT5G42710            |
| Brara.I01747 | F-box/leucine rich repeat protein                |
| Brara.I01748 | glucose-6-phosphate isomerase                   |
| Brara.I01749 | zinc finger FYVE domain containing protein       |
| Brara.I01750 | uncharacterized protein ath:AT5G42765            |
| Brara.I01751 | ZF-HD protein dimerization region               |
| Brara.I01752 | uncharacterized protein ath:AT5G42785            |
| Brara.I01753 | proteasome subunit alpha/beta                   |
| Brara.I01754 | dihydroflavonol 4-reductase                     |
| Brara.I01755 | inositol polyphosphate kinase 1                 |
| Brara.I01756 | U2 SNRNR auxiliary factor, small subunit         |
| Brara.I01757 | uncharacterized protein ath:AT5G42860            |
| Brara.I01758 | lipin 3-related                                  |
| Brara.I01759 | sterol carrier protein 2                        |
| Brara.I01760 | RNase_H                                          |

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Fig 3. Non-purple strains bear an insertion mutation in exon 4 of the *DFR* gene. (A) Diagram of predicted intron/exon structure of *DFR* with the insertion shown. Positions of primers used in this study are also shown. (B) PCR with primers pr7F and PR which are anchored in the 5th exon of the gene identifies an insertion mutation. (C) Nucleotide sequence of the insertion (GenBank accession KX379243). Flanking direct repeats are underlined. The two halves of the perfect palindrome are shaded. An in-frame stop codon is indicated with double underline. (D) PCR assay for wild type and mutant alleles of the *DFR* gene. In the top panel, PCR primers were prCF and prNR which amplify a product (N) in alleles with the insertion. In the bottom panel, PCR primers were prCF and prPR which amplify a product (P) from the wild type allele. In all reactions, primers pr1F and pr10R, which amplify a segment from a different part of the *DFR* gene (+), were

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AAAAAGGAATGAGAATTTCGCCAAAAAAACCTCAACCTTTACACGA
ATTGCCAAAAACAAACTGATACTTTGCCCCAAAAACACCA
AATTTCATTTAGACCTTTAGAATTTAATACATGTTTGGCTGACTT
GCCAACATTAGACCGCGTCACACAAAAATTTACAGAAATTTTACGT
CAGATTTTCTGTAAAATTGTTACGCCGTGCTCAAAATTGGCAAGT
CAGCCGAAAACATTGTAAAATTCTAAAGTCATTGAAAGTTTGT
GTTTTTTGAGCGGAGGCGACAAAATTCTGTTTTGGCGATTGTG
GTTGAGGTAGGATTTTTTTGGCGGATTTGG

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DFR Mutation in Fast Plants

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the mutant allele of the gene that can be used in an undergraduate or advanced high school teaching lab. Fig 5 shows the results of genotyping seedlings from stocks obtained from Carolina Biological Supply. The mutant and wild type alleles are readily detected in parental strains. In a sample of F2 generation seedling the PCR test shows that all nonpurple seedlings have the mutant allele only, and among the purple seedlings, some are have only the wild type allele and some carry the mutant allele in addition to wild type.

**Discussion**

Our data indicate that the *DFR* gene which encodes dihydroflavonol 4-reductase is the *anthocyaninless* gene of Rapid Cycling *Brassica rapa* (Fast Plants type). The non-purple trait maps to this position and the *DFR* gene on non-purple strains bears an insertion mutation that would doubtlessly inactivate the gene. If the palindromic sequence does not impede transcription and translation, it introduces a premature stop codon in the fourth exon of the gene. The linkage of loss of function mutation in *DFR* to the *anthocyaninless* locus is consistent with what is seen in other plant species. *DFR* encodes an enzyme that is essential for flavonol biosynthesis, and loss of function mutations have been found to result in a complete lack of anthocyanin pigments in maize [17], snapdragon [18], mouse ear cress [19], tomato [20], soybean [9], tobacco [21], Lewis’ monkeyflower [22], onion [23], Solonaceae [24], and Oriental lily [25].

Both nonpurple strains analyzed in this study have the insertion mutation. That is to be expected because although there are multiple nonpurple seed stocks of RCBr, they originate from the same original isolate (P. Williams, personal communication).

Educators at the advanced high school and introductory college level can employ a simple PCR assay to determine a plant’s genotype for *DFR* and relate this to a plant’s phenotype.

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**Table 3. Primers used to characterize the *DFR* gene.**

| Primer | Sequence               |
|--------|------------------------|
| pr1F   | CGAGCCAACGACATTCAT     |
| pr1R   | TCCACGTGACGTAAATCCT    |
| pr7F   | TAGCAGAAGAAGCAGCTTG    |
| pr10R  | TCTTGTCAGGTCTTTGCCT    |
| prCF   | CCAAGAAGATGACAGGTGG    |
| prNR   | TGGACGGCTGCTAAATTGG    |
| prPR   | TGGACCGATCAACATGTCG    |

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![Fig 4. Purple strain DWRCBr76 bears an insertion within the intron between exons 5 and 6 of the *DFR* gene (GenBank accession KX347549).](image)
Our work identifying the mutation responsible for the non-purple trait will be of great value for educators using Fast Plants for genetics instruction. Our findings are the first case of a trait in Fast Plants being characterized at the molecular level, and the recessive non-purple trait (genotype anl/anl) is the principle trait used in Mendelian genetics exercises developed by the Wisconsin Fast Plants Program [3]. Now the nonpurple trait is an observable phenotypic variation that can be used as a gateway to genomics. The genome of *Brassica rapa* has been sequenced in both Chinese cabbage [26] and in the FPSc strain of self-compatible Fast Plants [27]. Both can be searched and browsed in public databases. Students who study classical genetics with crosses of purple and nonpurple will now be able to relate this observable phenotypic variation to variation in a specific sequence and explore the genomic sequence data and browse the

![Fig 5. Development of a PCR test of the *anl* mutation for the teaching laboratory.](https://www.plosone.org/doi/10.1371/journal.pone.0161394.g005)

**Fig 5. Development of a PCR test of the *anl* mutation for the teaching laboratory.** (A) When DNA from a heterozygote is used as PCR template, wild type and mutant alleles cannot be simultaneously amplified. Gel contains products of duplicate reactions of Purple Stem, Hairy (PH), Nonpurple Stem, Yellow Green Leaf (NP), and a hybrid of the two (F1). (B) A PCR test that detects the mutant (N) and wild type (P) alleles in separate reactions in plants grown from Carolina Biological Supply seeds. PCR is as described in Fig 4.D. Lanes: (1) DNA ladder, (2) Nonpurple Stem, Yellow Green Leaf, (3) Purple Hairy, (4) F1 generation, (5–8) F2 seedlings with nonpurple phenotype, (9–20) F2 seedlings with purple phenotype.

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chromosome around the DFR gene. It is also a gateway to connecting an easily observable trait, purple stem color, with a major biochemical pathway, flavonoid biosynthesis.

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Author Contributions
Conceived and designed the experiments: DLW.
Performed the experiments: DLW AV GS.
Analyzed the data: DLW AV GS.
Contributed reagents/materials/analysis tools: DLW.
Wrote the paper: DLW.

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