

A Single SNP in an Evolutionary Conserved Region within Intron 86 of the *HERC2* Gene Determines Human Blue-Brown Eye Color

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We have previously demonstrated that haplotypes of three single nucleotide polymorphisms (SNPs) within the first intron of the *OCA2* gene are extremely strongly associated with variation in human eye color. In the present work, we describe additional fine association mapping of eye color SNPs in the intergenic region upstream of *OCA2* and within the neighboring *HERC2* (ect domain and RLD2) gene. We screened an additional 92 SNPs in 300–3000 European individuals and found that a single SNP in intron 86 of *HERC2*, rs12913832, predicted eye color significantly better (ordinal logistic regression $R^2 = 0.68$, association $LOD = 444$) than our previous best *OCA2* haplotype. Comparison of sequence alignments of multiple species showed that this SNP lies in the center of a short highly conserved sequence and that the blue-eye-associated allele (frequency 78%) breaks up this conserved sequence, part of which forms a consensus binding site for the helicase-like transcription factor (HLTF). We were also able to demonstrate the *OCA2* R419Q, rs1800407, coding SNP acts as a penetrance modifier of this new *HERC2* SNP for eye color, and somewhat independently, of melanoma risk. We conclude that the conserved region around rs12913832 represents a regulatory region controlling constitutive expression of *OCA2* and that the C allele at rs12913832 leads to decreased expression of *OCA2*, particularly within iris melanocytes, which we postulate to be the ultimate cause of blue eye color.

Introduction

Human eye color is a polymorphic phenotype under strong genetic control.¹ The gene responsible for oculocutaneous albinism type II (*OCA2*)^{2–4} has hitherto seemed the best candidate to explain the genetic linkage of blue-brown eye (BEY2/EYCL3 [MIM 227220]) and brown hair (HCL3 [MIM 601800]) color to chromosome 15q11.2–q12.^{5–11} *OCA2* is the human homolog of the mouse pink-eyed dilution gene (*p*).¹² The human *OCA2* gene is divided into 24 exons covering >345 kbp of DNA; 23 of these exons span the 836 amino acid coding region, with exon 1 representing exclusively a noncoding 5'UTR.¹³ The resulting gene product, the P protein, is an integral membrane protein containing 12 transmembrane spanning regions that helps regulate melanogenesis.¹⁴

Many polymorphisms in *OCA2* occur in different populations with at least 13 nonsynonymous amino acid substitutions reported.^{2,13,15–18} Only two of these were found to be present at significant frequency in our previous studies of adolescent twins and family members of mainly European descent from southeast Queensland, Arg305Trp (rs1800401) and Arg419Gln (rs1800407) at 0.05 and 0.09, respectively.¹⁸ They exhibited only a minor impact on eye color, leading us to conclude that *OCA2* coding alleles account for only a small proportion of the variation in iris pigmentation, at least within fair-skinned populations.

We proceeded therefore to screen tagging SNPs throughout the entire *OCA2* locus. The highest association for blue:nonblue (green/hazel or brown) eye color was found

with three SNPs in intron 1: rs7495174 T/C, rs6497268 (now rs4778241) G/T, and rs11855019 (now rs4778138) T/C. We found the TGT/TGT diplotype in 62.2% of samples, and this was the major blue-eye genotype, with a frequency of 91% in blue- or green-eyed individuals, compared with only 9.5% in those with brown eyes.¹⁸ The position of this major diagnostic haplotype for eye color suggested that differences within the 5' proximal regulatory control region of *OCA2* alter temporospatial expression of the gene and may be responsible for these associations.

To further refine the elements controlling eye color, we have tested for association with haplotype-tagging SNPs proximal to intron 1 of *OCA2*, which span the intergenic region and encompass the 3' end of the upstream gene, *HERC2*. We describe the fine mapping of SNPs that are better predictors of blue-brown eye color than the existing *OCA2* intron 1 TGT haplotype block. Moreover, we propose a mechanism whereby a single base change (our best predictive SNP), contained within a highly conserved region of the *HERC2* gene intron 86, may abrogate accessibility of the *OCA2* chromosomal region by transcription factors necessary for expression of *OCA2* in people of different eye color.

Material and Methods

Structure of the Study Population and Pigmentation Characteristics

Adolescent twins and their siblings were recruited for an investigation of genetic and environmental factors contributing to the

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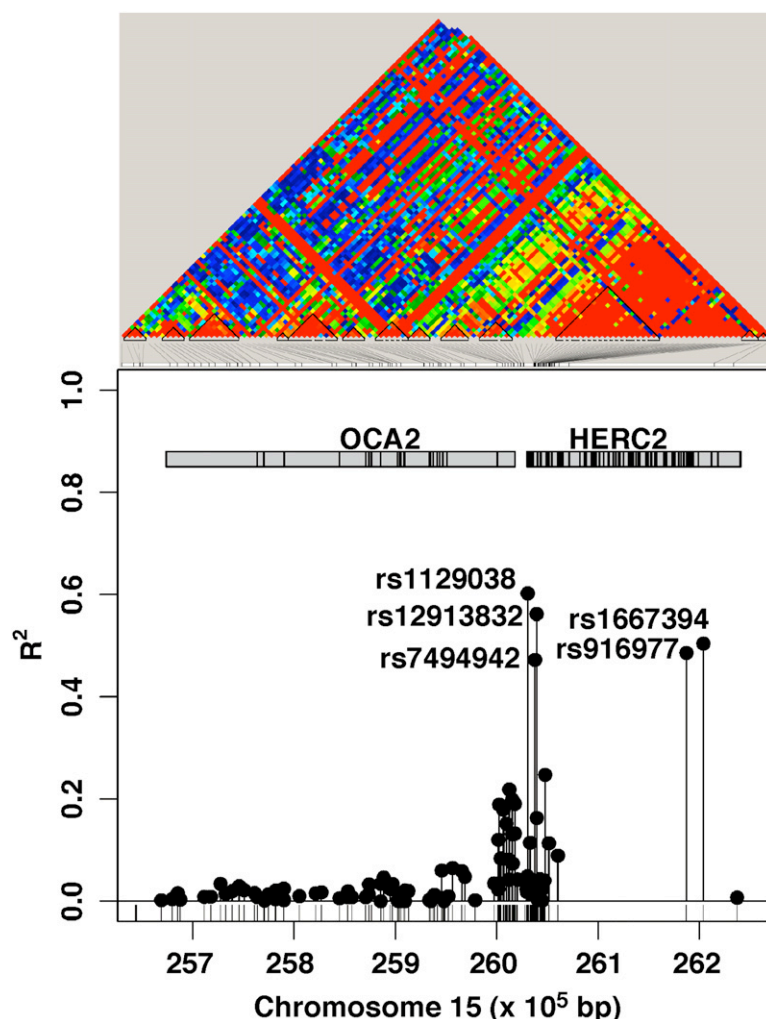


Figure 1. Association of *OCA2* and *HERC2* SNPs with Eye Color

Association of *OCA2* and *HERC2* SNPs with eye color (three-point scale) measured as R^2 in ordinal logistic regression analysis in the lower panel. The upper panel shows the linkage disequilibrium (r^2) between the SNPs.

OCA2 and *HERC2* SNP Genotyping

Genotyping at a set of 58 *OCA2* SNPs was described previously.¹⁸ We designed assays for another 92 SNPs including 30 within *OCA2*, 7 SNPs within the intergenic region, and 55 within *HERC2*, concentrating on 3' introns/exons. These were selected based on their location within the gene and/or differences in published allele frequencies among the three HapMap populations. They were designed to be assayed in three multiplexes, so SNPs were typed with iPLEX chemistry on a Compact MALDI-TOF Mass Spectrometer (Sequenom, Inc., San Diego, CA) by standard methods.²⁴ In a subset of twins, we also utilized SNPs genotyped as part of a genome-wide association study with the Affymetrix 100K array.

Statistical Methods

We performed standard linear and logistic regression analyses of the data with the R computer package (Version 2.5, R Development Core Team 2007). We used SOLAR 4.0.7 and MENDEL 7.01 to perform individual SNP and haplotypic association analysis correctly allowing for the relatedness within the sample. Some simulation-based p values for association have been generated with the Sib-pair program.¹⁸ Long-range haplotypes were inferred with Beagle²⁵ and Haploview 4.²⁶

Results

Association of *OCA2-HERC2* SNP Alleles and Haplotypes with Blue Eye Color in an Adolescent Twin Collection

The eye color grade distributions were similar to our earlier reports,^{18,27} with 46.1% blue/gray, 27.7% green/hazel, and 26.3% brown in the total sample collection. The genotyped subjects were approximately 52% female and 48% male and there were no significant gender differences in eye color distribution.

In our first round of follow-up genotyping of 8 SNPs 5' of the *OCA2* gene, the most strongly associated of the SNPs was rs12913832 ($p = 2 \times 10^{-78}$). Surprisingly, this SNP was 21.1 kb upstream of the *OCA2* first exon, in intron 86 of the *HERC2* gene (Figure 1). Individuals carrying the C/C genotype had only a 1% probability of having brown eyes. By contrast, T/T carriers had an 80% probability of being brown eyed.

We were able to show that this SNP gave additional information over that obtained with our previous best

development of pigmented nevi and were also phenotyped for pigment traits including skin, hair, and eye color. The pigmentation characteristics of the twins were examined on up to three occasions, at 12, 14, and 16 years of age as previously described.¹⁹ Subjects were overwhelmingly (>95%) of northern European origin (mainly Anglo-Celtic). Eye color was rated by one research nurse (AE) as blue/gray (1), green/hazel (2), or brown (3) and cross-validated with the individual's self-report.¹⁹

There were 5075 family members in 1100 pedigrees with some phenotypic data, and DNA was available for genotyping for 3839 individuals within 1037 of these pedigrees. Excluding one member of each genotyped MZ twin pair, there were 3011 individuals with complete *OCA2* genotype, hair color, eye color, and sex recorded. We analyzed the data collected when the twins were 12 years old, because these are most complete.

We also carried out genotyping of an additional collection of Queensland cutaneous malignant melanoma (CMM) cases and their family members. In brief, the Queensland Familial Melanoma Project ascertained all 10407 CMM cases diagnosed from 1982 to 1990 in the state of Queensland, and a sample of families stratified on strength of family history was drawn.²⁰⁻²³ Phenotypes including hair, skin, and eye color were assessed by questionnaire, and DNA from available cases and selected unaffected members was collected. Only a subset of 96 blue-eyed and 96 brown-eyed individuals from this sample was genotyped for the present study.

Table 1. Haplotype Frequencies at *HERC2*-*OCA2* Locus SNPs rs12913832-rs7495174-rs6497268-rs11855019 versus Eye Color

Haplotype	Total	Blue	Green-Hazel	Brown
C-TGT	0.73547	0.91268	0.74083	0.37359
C-TTC	0.02627	0.02661	0.02115	0.02377
C-TGC	0.01031	0.01458	0.00260	0.01140
C-TTT	0.00768	0.01447	0.00345	0.0
C-CTC	0.00202	0.00115	0.00249	0.00004
C-CGT	0.00120	0.00131	0.00054	0.00328
C-CGC	0.00021	0.0	0.0	0.00204
T-TTT	0.07598	0.01260	0.11152	0.15577
T-CTC	0.05431	0.00708	0.03599	0.16711
T-TGT	0.05224	0.00318	0.05441	0.15851
T-TGC	0.02482	0.00393	0.02309	0.07032
T-TTC	0.00830	0.00212	0.00298	0.03270
T-CGT	0.00059	0.0	0.00096	0.0
T-CTT	0.00049	0.00030	0.0	0.00146
T-CGC	0.00011	0.0	0.0	0.0

predictor, the three SNP haplotype in the first intron of *OCA2*. A haplotype analysis combining the original three SNPs with rs12913832 showed that the latter “split” the original haplotypes more precisely into eye color groups (see Table 1). Most critically, the C-TGT haplotype has a frequency of 91% in blue-eyed individuals and 37% in brown-eyed individuals, whereas the T-TGT haplotype is present in 0.3% of blue-eyed individuals and 15% of brown-eyed individuals (odds ratio = 122). Confirming this, a multiple ordinal logistic regression including these four SNPs found that the original three SNPs could be dropped from the model without a significant loss of predictive power.

Assay of *HERC2* SNP Alleles and Haplotype Association in Individuals Selected for Blue or Brown Eye Color

We then carried out genotyping at an additional 92 SNPs within the *OCA2* and *HERC2* genes in a subsample of 384 individuals from our collection (192 with blue eyes, 192 with brown eyes), choosing only one person from any family. Only one SNP, rs1129038, was found to slightly more accurately predict eye color than rs12913832 in this small subsample (Figure 1). This was located in the 3'UTR of the *HERC2* gene exon 93 and 12.4 kb upstream of the *OCA2* first exon. This had similar genotype frequencies to rs12913832 among blue-eyed and brown-eyed individ-

uals. In addition, two more distant SNPs exhibited similar though slightly smaller effects on eye color: rs1667394 and rs916977, in introns 4 and 12 of *HERC2*, respectively. Further examination of these four eye color SNPs confirmed that they were all in strong linkage disequilibrium with one another (see Table 2). However, it was possible to show that rs1129038 and rs12913832 were more likely to be causative or in strong LD with the causative variant (see Table 3). Specifically, the rs1129038-rs12913832-rs916977-rs1667394 **ACGA/ACGA** genotype had a blue-eye phenotype whereas the **ACGA/GTGA** genotypes were brown-eyed ($p = 2 \times 10^{-16}$).

Assay of *HERC2* SNP Alleles and Haplotype Association with Eye Color in an Adolescent Twin Collection

We used the Beagle program²⁵ to impute the most likely haplotypes for our sample of 384 individuals including all genotyped SNPs between R419Q in *OCA2* and rs1667394 in *HERC2* (129 SNPs). When we tested for an optimum combination of alleles from these haplotypes to best predict eye color by a recursive partitioning method,²⁸ rs1129038 and rs12913832 were selected (Bonferroni-corrected $p < 10^{-48}$) (Figure 2), along with R419Q (included at a far lower level of statistical significance, $p = 0.041$).

Based on these results, we elected to genotype only these two SNPs in our entire collection. The total number of individuals genotyped at these key SNPs was 3961. In the total sample, these two SNPs were confirmed to be in almost perfect LD. Of the rs1129038-rs12913832 **AC** haplotype homozygotes, 99% were nonbrown (blue or green/hazel) in eye color, and 99% of **GT** haplotype homozygotes were nonblue in eye color (Table 4). These results closely match those expected for the brown eye color *BEY2* locus based on segregation analysis. It was not possible by statistical methods to further decide between these two SNPs as the best candidate for a causal variant.

We then used multivariate approaches to further dissect effects of other variants within *OCA2* via the complete data set. We were able to confirm that the *OCA2* R419Q substitution is independently associated with eye color. The most striking effect was seen on the rs12913832*T/T background, where the penetrance for green/hazel eyes for 419Q/419Q, 419Q/419R, and 419R/419R was then 50%, 21%, and 6%, respectively ($p = 0.0003$). In a parallel analysis, we examined the effects of these two SNPs on risk of

Table 2. Minor Allele Frequencies and Measures of Intermarker Linkage Disequilibrium for the Four *HERC2* Locus SNPs Most Strongly Associated with Eye Color in Our Sample

SNP	Sequence Position (bp)	MAF	rs1129038	rs12913832	rs916977	rs1667394
rs1129038	26,030,454	0.236 (G > A)	1	0.984	0.556	0.588
rs12913832	26,039,213	0.220 (C > T)	0.997	1	0.546	0.580
rs916977	26,186,959	0.204 (G > A)	0.989	0.978	1	0.928
rs1667394	26,203,777	0.214 (A > G)	0.990	0.979	1	1

MAF, minor allele frequencies; r^2 above diagonal; D' below diagonal.

Table 3. Phased Genotypes at Four Key *HERC2* Locus SNPs versus Eye Color

	rs1129038	rs12913832	rs916977	rs1667394	Phased ^a	Blue	Brown
1 ^b	A/A	C/C	G/G	A/A	ACGA/ACGA	169	1
2 ^{b,c,d}	A/G	C/T	G/G	A/A	ACGA/GTGA	1	47
3 ^{c,d}	A/G	C/T	A/G	A/G	ACGA/GTAG	10	81
4 ^d	A/G	C/T	G/G	A/G	ACGA/GTGG	0	8
5	G/G	T/T	A/A	G/G	GTAG/GTAG	0	18
6	G/G	T/T	A/G	A/G	GTAG/GTGA	0	14
7	A/A	C/C	A/G	A/G	ACGA/ACAG	1	1
8	A/G	C/C	A/G	A/G	ACGA/GCAG	1	1
9	A/G	C/C	G/G	A/A	ACGA/GCGA	1	1
10	G/G	T/T	A/G	G/G	GTAG/GTGG	0	1
11	G/G	T/T	G/G	A/A	GTGA/GTGA	0	3

^a **AC** phased rs1129038 and 12913832 in bold.

^b The Fisher exact test comparing counts for row 1 versus 2, $p = 2 \times 10^{-16}$.

^c Row 2 versus 3, $p = 0.097$.

^d Row 2 versus 3 versus 4, $p = 0.195$.

cutaneous malignant melanoma. Interestingly, R419Q was a significant independent risk factor (allelic odds ratio adjusted for the other locus, OR = 1.27, empirical $p = 0.0006$), as previously found by ourselves and others, and rs12913832 was less strongly associated (OR = 1.15, $p = 0.008$).

Sequence Alignment of *HERC2* Exon 86 to Exon 93 across Species

We examined cross-species similarity in alignment of the 10.4 kb human *HERC2* genomic sequence from exon 86 to exon 93 (GenBank NT_010280.17), encompassing the interval between our two peak eye color SNPs. Alignments were first tested by comparing each individual exonic and intronic segment separately with the corresponding mouse *Herc2* genomic sequence (GenBank NT_039424.7). Apart from the exonic coding regions, there was only one region of marked similarity found—between the human and mouse intron 86 sequences, as seen in a dot matrix plot (Figure 3A). The GenBank reference sequence for *HERC2* was also compared with the Celera sequence (Celera NW_925783) across this 10.4 kb region, and 23 nucleotide differences were apparent in this alignment, notably the rs1129038-rs12913832 SNP haplotype was the brown

GT in GenBank and blue AC in the Celera sequences, respectively, consistent with the eye color of one of the individuals used to generate the Celera database.²⁹ The alignment of the conserved region of intron 86, between the two versions of the human 406 bp and corresponding 420 bp mouse sequence, showed more than 77% shared identity (Figure 3B). An additional SNP rs6497271 contained within this region was also identified. To search for additional variants in the immediate vicinity of our key SNPs, we sequenced the regions around rs12913832 and rs1129038 in five individuals with brown eyes and five individuals with blue eyes (690 bp either side of rs12913832 and 600 bp either side of rs1129038; primers available on request). No novel variants were found in these regions.

An alignment of this interval across multiple species, concentrating especially on the regions close to our two peak SNPs, was also examined with the UCSC browser Multiz alignment track³⁰ and in the Vista Enhancer “Computational Dataset.”³¹ The latter resource flagged only two regions of *HERC2* as strongly conserved and thus likely to contain enhancer elements—portions of intron 2 (which contains no known SNPs) and intron 86 (containing rs12913832 and rs6497271). Although we observed little

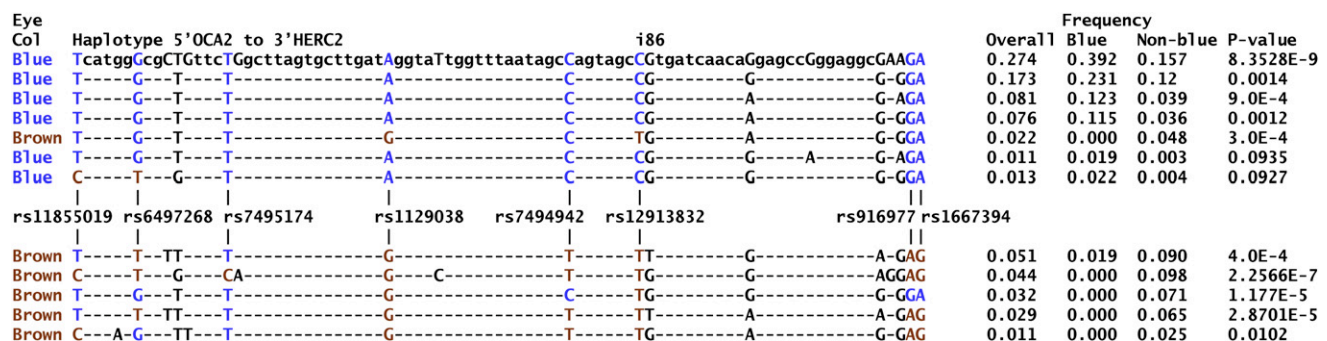


Figure 2. SNP Haplotypes Associated with Eye Color

SNP haplotypes associated with blue and nonblue eye color spanning rs11855019 to rs1667394 of the OCA2-HERC2 locus. The SNP bases associated with blue and brown haplotypes are indicated by color.

Table 4. Genotypes with Penetrances Given in Parentheses for *HERC2* rs12913832 and *OCA2* R419Q versus Eye Color in the Complete Sample Collection

rs12913832	R419Q	Blue	Green	Brown
C/C	R/R	1268 (0.71)	499 (0.28)	13 (0.01)
C/C	R/Q	125 (0.83)	25 (0.17)	1 (0.01)
C/C	Q/Q	1 (1.00)	0 (0.00)	0 (0.00)
C/T	R/R	38 (0.05)	214 (0.29)	496 (0.66)
C/T	R/Q	36 (0.12)	176 (0.58)	91 (0.30)
C/T	Q/Q	0 (0.00)	9 (0.75)	3 (0.25)
T/T	R/R	2 (0.02)	7 (0.06)	100 (0.92)
T/T	R/Q	0 (0.00)	12 (0.22)	42 (0.78)
T/T	Q/Q	0 (0.00)	5 (0.50)	5 (0.50)

sequence conservation around rs1129038, there was again striking conservation immediately around rs12913832 (Figure 3C). The Regulatory Potential mammalian conservation scoring method provided by the ESPERR 7 program³² gave a score 10-fold higher than the baseline signal (a level found to predict enhancer activity in 60%–100% of cases when validated by chromatin immunoprecipitation or integration of the putative expression cassette via the *Cre-loxP* system³³).

Analysis of this sequence with the MatInspector Program³⁴ suggests that this may represent a HLTF (SMARCA3) binding site. Tellingly, this consensus sequence is abolished in the rs12913832*C blue-eyed allele. Other potential transcription factor binding sites relevant to melanocytic cell gene expression identified in this search are two strong *MITF* sites³⁵ and three *LEF1* sites. Notably, rs6497271 is in the third of these *LEF1* sites, and presence of the minor allele results in the failure of this motif to be identified in a MatInspector search. rs6497271 had only a 1% minor allele frequency (MAF) within our sample, so it cannot explain the association to the region.

Discussion

There is overwhelming evidence implicating the *OCA2* gene region in regulation of human pigmentation. Mutations in the *OCA2* gene lead to oculocutaneous albinism. Deletion of the region encompassing the *OCA2* gene on chromosome 15 as observed in Prader-Willi and Angelman syndromes is associated with hypopigmentation of the skin, hair, and eyes, and extra copies of this chromosomal region result in generalized hyperpigmentation of the skin. Normal variation in eye color shows strong genetic linkage⁸ and association¹⁸ to markers in the *OCA2* locus. In our previous combined segregation-linkage analysis in this twin sample, we estimated the frequencies of a dominant brown eye *B* allele as 21% and recessive *b* allele as 79%, which was close to the 26% *B* and 74% *b* allele modeled for the US white population by Hasstedt.³⁶ The allele frequencies and penetrances estimated for rs12913832 in our Anglo-Celtic population match these predictions almost exactly. The European population frequencies in

Hapmap for this SNP are in agreement (MAF 21%), and the variant is absent from other population groups.³⁷ There is strong and extensive linkage disequilibrium across the *OCA2-HERC2* region, as well as marked differences in SNP allele frequencies between different ethnic groups. Voight et al.³⁸ used HapMap data to show that this can be interpreted as a signal of positive genetic selection in Europeans, similar to that seen around other pigmentation loci such as *TYRP1*, *DTNBP1*, and *SLC24A5*. The hitchhiking seen around such loci can decrease the power of association-based fine mapping.

Although rs12913832 lies within a distal intron of *HERC2* (intron 86 of a 93 exon encoded gene spanning 211 kb), we do not believe that the *HERC2* protein is involved in regulating the pigmentation pathway. Three mouse deletions that map within this distal region of *HERC2*, which do not include the *OCA2* locus, were originally labeled as *p* gene mutants because they exhibit pigmentation changes that are intermediate to those seen in canonical *p* mutant homozygotes: dark pink eyes with mottling or partially decreased eumelanin quantities in hair.^{39–41} Effects on pigmentation are thought to result from sequences within this region of *HERC2* controlling expression of *OCA2*. One of the radiation-induced mutant mouse alleles characterized by Lehman et al.,⁴⁰ *p*^{bs} (black eyed, sterile; originally *p*^{24H42}), involves a deletion of only 8 kb in *Herc2*, which spans the orthologous region to human *HERC2* exons 86–93. Lehman et al. comment in parentheses that “[t]he relatively mild hypopigmentation associated with the *p*^{bs} deletion apparently reflects an effect of the deletion on expression of the closely linked *p* gene.” Walkowicz et al.⁴¹ describe characterization of the *p*^{12DTR}, *p*^{103G}, and *p*^{39DSD} alleles.⁴³ These all involve deletions or rearrangements of *Herc2*. Those authors concluded that elements regulating the expression of *p* probably lie within *Herc2*.

Recently, strong association between eye color and rs1667394 in *HERC2* was reported in Icelandic population,⁴⁴ but we excluded this as the casual SNP based on haplotype analysis. One of our two most strongly associated SNPs, rs12913832, lies within an evolutionarily conserved 406 bp region. Moreover, database searches of this region revealed transcription factor binding sites for *LEF1*, *HLTF*, and *MITF*. Both *LEF1* and *MITF* are critically important to gene regulation in melanocyte cell development, differentiation, and tissue-specific transcription.^{45,46} Helicase-Like Transcription Factor (HLTF) is a member of the *SWI2/SNF2* family, DNA-dependent chromatin-remodelling ATPases that have been implicated in a wide variety of processes involving the modification of chromatin configuration to allow access of the transcriptional machinery. HLTF is the only member of the family in humans that incorporates a specific DNA recognition site that binds a variety of gene promoters, including *PAI-1* and beta-globins (where it seems to be involved in controlling levels of constitutive expression),^{47,48} uteroglobin (transducing of the effects of prolactin) on

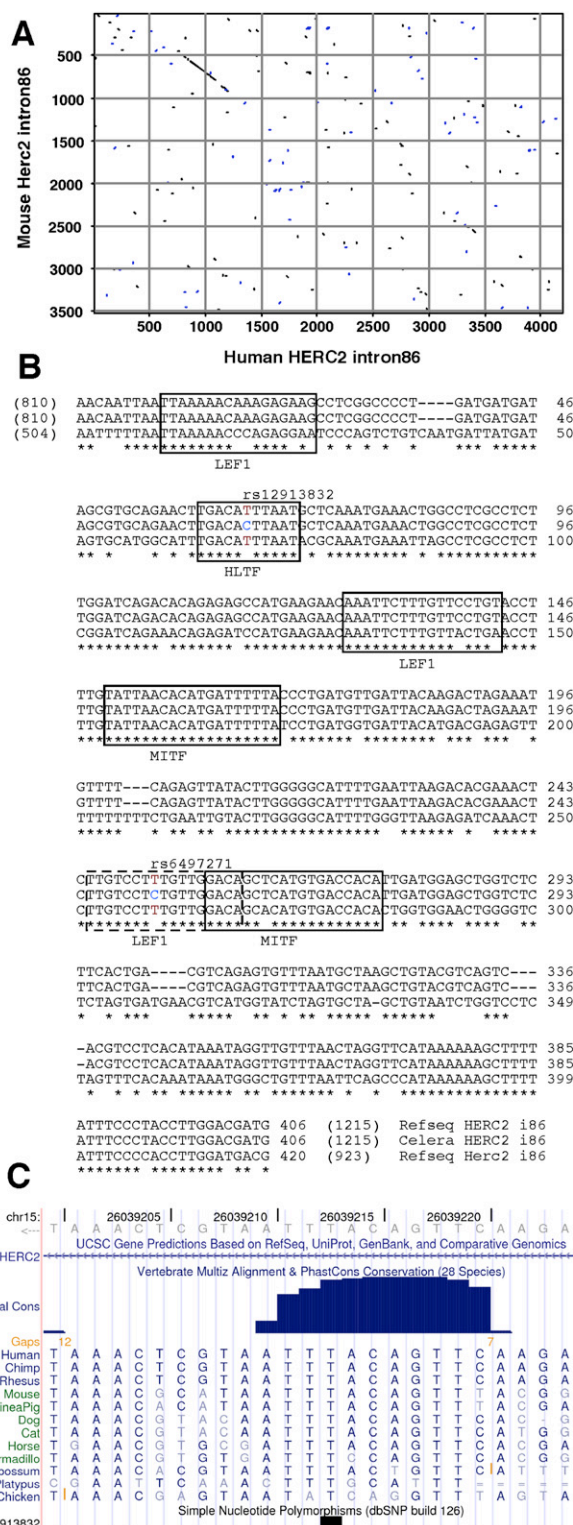


Figure 3. Conserved Sequences in *HERC* Intron 86

(A) Dot-matrix-aligned sequence comparison of intron 86 from human *HERC2* (x axis) and mouse *Herc2* (y axis) with the MacVector program (MacVector, Inc.).

(B) ClustalW alignment of the sequence similarity found in (A) with the RefSeq and Celera human with the RefSeq mouse sequences. Transcription factor binding sites identified by the MatInspector Program are boxed with SNP changes shown in color.

uteroglobin,⁴⁹ and an enhancer region of the myosin light chain locus.^{50,51} The binding site in intron 86 matches the consensus recognized by the HLTF variant known as RUSH-1- α as determined by CASTing.⁴⁹

Another SWI/SNF family member, BRG1, has recently been shown to be involved in pigmentation pathways, being recruited by MITF to melanocyte-specific promoter regions to induce changes in chromatin structure at endogenous loci, and thus initiating the process of melanocyte differentiation.⁵² We therefore suggest that the presence of both MITF and HLTF enhancer sequence elements in the evolutionary conserved region of *HERC2* intron 86 is a further pointer that this is a locus control region that determines the expression of the *OCA2* gene product.

We have now genotyped most of the reported SNPs in the 3' end of *HERC2* and the 5' end of *OCA2*, finding a large number (39 out of 93 or 42% in the second round of genotyping) to be monomorphic in our sample. The blue-eye-color-associated allele must be reasonably common if a single SNP is to explain most of the variation in eye color in European-descended populations. This makes it less likely that another SNP as yet unidentified in the region is the true causative variant. Based on the foregoing, we conclude that the conserved region around rs12913832 represents a regulatory region controlling constitutive expression of *OCA2*, and that the C allele at rs12913832 leads to decreased expression of *OCA2*, particularly within iris melanocytes. We speculate that the regulatory mechanism is abrogation of the binding site for HLTF that regulates transcription of the neighboring *OCA2* gene. We also confirmed that the common coding variant *OCA2**R419Q acts to modify the penetrance of this locus and that this effect includes modification of the risk of malignant melanoma.

Acknowledgments

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(C) UCSC Genome Browser⁵³ screen shot showing sequence alignment for seven species and associated phastCONS scores for sequence around rs12913832 of *HERC2* intron 86. Sequence is shown in the 3' to 5' direction of the upper strand.

Web Resources

The URLs for data presented herein are as follows:

dbSNP, <http://www.ncbi.nlm.nih.gov/SNP/>

HapMap, <http://www.hapmap.org/>

Online Mendelian Inheritance in Man (OMIM), <http://www.ncbi.nlm.nih.gov/Omim>

UCSC Genome Browser, <http://genome.ucsc.edu/>

VISTA Enhancer Browser, <http://enhancer.lbl.gov/>

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