



Título artículo / Títol article: Gender is a major factor explaining

> discrepancies in eye colour prediction based on HERC2/OCA2 genotype and the

IrisPlex model

Autores / Autors Conrado Martínez-Cadenas, María Peña-

Chilet, Maider Ibarrola-Villava, Gloria Ribas

Revista: Forensic Science International: Genetics

Versión / Versió: Versió post-print de l'autor

Cita bibliográfica / Cita

bibliogràfica (ISO 690): MARTINEZ-CADENAS, Conrado, et al.

> Gender is a major factor explaining discrepancies in eye colour prediction based on< i> HERC2/OCA2</i> genotype and the IrisPlex model. Forensic Science International: Genetics, 2013, vol. 7, no 4,

p. 453-460.

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Gender is a major factor explaining discrepancies in eye colour prediction based on HERC2/OCA2 genotype and the IrisPlex model

QI, Conrado Martinez-Cadenas, a, Maria Peña-Chilet, b, Maider Ibarrola-Villava, b, Gloria Ribas b,*

ARTICLE INFO

Article history: Received 23 January 2013 Received in revised form 13 March 2013 Accepted 17 March 2013

Keywords: Genetics Pigmentation Eve colour Irisplex Polymorphisms Spanish population

ABSTRACT

In recent years, several studies have greatly increased our understanding of the genetic basis underlying human eye colour variation. A large percentage of the eye colour diversity present in humans can already be genetically explained, so much so that different DNA-based eye colour prediction models, such as IrisPlex, have been recently developed for forensic purposes. Though these models are already highly accurate, they are by no means perfect, with many genotype-phenotype discrepancies still remaining unresolved. In this work we have genotyped six SNPs associated with eye colour (IrisPlex) in 535 individuals from Spain, a Mediterranean population. Aside from different SNP frequencies in Spain compared to Northern Europe, the results for eye colour prediction are quite similar to other studies. However, we have found an association between gender and eye colour prediction. When comparing similar eye colour genetic profiles, females tend, as a whole, to have darker eyes than males (and, conversely, males lighter than females). These results are also corroborated by the revision and metaanalysis of data from previously published eye colour genetic studies in several Caucasian populations, which significantly support the fact that males are more likely to have blue eyes than females, while females tend to show higher frequencies of green and brown eyes than males. This significant gender difference would suggest that there is an as yet unidentified gender-related factor contributing to human eye colour variation.

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1. Introduction

Eye colour variability is one of the most visible manifestations of individual physical variation in people of Caucasian descent. The pigmentation of the iris is determined by the amount of melanin present in the anterior layer of the iris stroma [1]. Eye colour is thus a quantitative character dependent on melanin content, with a gradient of different colours ranging from light blue (very low melanin content) to dark brown (high melanin content). However, in almost all cases this quantitative variability can be unequivocally reduced to three broad qualitative categories naturally perceived by the human eye: blue/grey, green/hazel (also called intermediate) and brown/black.

1872-4973/\$ - see front matter © 2013 Published by Elsevier Ireland Ltd. http://dx.doi.org/10.1016/j.fsigen.2013.03.007

Human eye colour is a fully genetically determined trait with a complex inheritance pattern, although most times it apparently behaves as a simple Mendelian character. This is so because one genetic region, HERC2/OCA2, accounts for the majority of the blue and brown variation (the two most common colours) in human eve colour [2,3].

Although genetic association studies had already identified the relationship between the OCA2 gene and eye colour [2,4], it was not until 2008 when three independent studies found that HERC2, a neighbouring gene, and specifically the SNP rs12913832, was the key human eye colour regulator [5-7]. In addition, other genes such as SLC24A4, SLC45A2, TYR, TYRP1, ASIP or IRF4 have also been recognised to contribute to eye colour variation, although to a much lesser extent [8-10].

More recently, investigations have focused on the development of a forensic genetic test to accurately predict eye colour from an unknown or anonymous DNA sample [11-13]. Although these studies provide slightly different recommendations for accurate eye colour prediction, all suggest that the genotyping of a small number of SNPs from a handful of pigmentation-associated genes can predict blue/brown eye colour with more than 90% accuracy.

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^a Department of Medicine, University of Castellon Jaume I, Castellon, Spain

^b Medical Oncology and Haematology Unit, Health Research Institute – INCLIVA, Valencia, Spain

Corresponding author at: Medical Oncology and Haematology Unit, Health Research Institute – INCLIVA, Av. Blasco Ibáñez, 17, 46010 Valencia, Spain Геl.: +34 963862894.

E-mail address: gribasdespuig@gmail.com (G. Ribas).

¹ These authors contributed equally to this work.

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Significantly, intermediate eye colours such as green or hazel still remain quite difficult to predict [8].

In this work, as in Walsh et al. [13], we have genotyped six SNPs associated with eye colour (IrisPlex) in 535 people from Spain, a Mediterranean population. Aside from different SNP frequencies in Southern European populations when compared to Northern European ones, results for eye colour prediction are similar to other studies. However, we have found a very strong association between gender and eve colour prediction. When comparing similar eye colour genetic profiles, females tend, as a whole, to have darker eyes than males (and, conversely, males lighter than females). This significant gender difference would suggest that there is an as yet unidentified factor contributing to human eye colour variation.

2. Materials and methods

2.1. Study subjects and data collection

A total of 535 individuals were included in this study. All samples belonged to healthy control subjects of Spanish origin drawn from various skin cancer studies, and were collected at several Spanish hospitals from the provinces of Madrid, Valencia and Castellon (Supplementary Table S1). All participants provided written informed consent, and the study was approved by the Ethics Committee of INCLIVA Health Research Institute, Valencia, Spain, Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.fsigen.2013.03.007.

A standardised questionnaire was used to collect information on pigmentation characteristics, as well as personal and family history of relevant diseases. For the purpose of this study we took into consideration age, gender and eye colour. Eye colour was determined by especially trained personnel. All eye colours were grouped in just three functional categories: brown/black, hazel/ green (or intermediate), and blue/grey.

Genomic DNA from individuals was isolated from peripheral blood lymphocytes and diluted to a final solution of 50 ng/ μ l. This was done using the traditional saline method or the DNAzol procedure (Invitrogen, Eugene, OR, USA). DNA concentration was quantified in samples prior to genotyping by using Quant-iT PicoGreen dsDNA Reagent (Invitrogen, Eugene, OR, USA). Further DNA concentration measurements were obtained using the Nanodrop 2000 spectrophotometer.

2.2. SNP selection

We genotyped the six SNPs included in the IrisPlex eye colour panel [8,13]. These comprise rs12913832 (located in the HERC2/OCA2 region), rs1800407 (OCA2 gene), rs12896399 (SLC24A4 gene), rs16891982 (SLC45A2 gene), rs1393350 (TYR gene), and rs12203592 (IRF4 gene). These six SNPs have been used in previous studies, and have already demonstrated their suitability for predicting eye colour in Caucasian populations [12–16]. Public databases such as NCBI (http://www.ncbi.nlm. nih.gov) and Ensembl (http://www.ensembl.org) were used to collect information about SNPs and genes. A north European population (CEU) and a southern one from Tuscany, Italy (TSI) available from the HapMap project (phase 3), were used to get all SNP minor allele frequencies, as well as the raw genotypes. Details of gene names, gene locations, nucleotide changes, minor allele frequencies, SNP assay number and eye colour associated with the less frequent allele are provided in Table 1 and in Supplementary Table S2. Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.fsigen. 2013.03.007.

2.3. Genotyping assays

Genotyping assays were carried out using Kaspar technology (KBiosciences, Hoddesdon, UK). PCRs were performed in a total reaction volume of 4 µl containing about 10 ng of genomic DNA, a final concentration of 4× New Kaspar Reaction Mix and 12 μm of each Kaspar primer. For SLC45A2 SNP rs16891982, TaqMan technology was used (Applied Biosystems, Foster City, USA).

PCR conditions varied depending on the requirements of each probe, always according to the manufacturer's indications. The genotype of each sample was determined by measuring final allele-specific fluorescence in the ABI Prism 7900HT Detection System, using SDS v2.3 software for allelic discrimination (Applied Biosystems, Foster City, USA).

As a quality control measure, we included one no-template sample and one sample duplicate in each 96-well plate (a total of four per 384-well plate used). Genotypes were provided automatically by the software and were confirmed manually in the laboratory by two different personnel. Additionally, all apparent discrepancies between genotype and phenotype were genotyped twice in independent experiments in order to prevent potential genotyping errors. Only individuals without missing genotypes (493) were considered for further analyses.

2.4. Prediction model

Liu and colleagues [8] have previously published the IrisPlex algorithm for eye colour prediction used in this and previous studies [13]. It is based on a multinomial logistic regression model. The probabilities of each individual for being brown-eyed (π_1) , blue-eyed (π_2) , and otherwise (π_3) were calculated based on the

Q5 Allelic distributions in different European populations.

Gene	SNP	Minor allele	Minor allele's colour prediction	Ancestral allele	MAF HapMap_CEU	MAF HapMap_TSI	Spanish populati	on	p-Value CEU	p-Value TSI
							pHWE	MAF		
HERC2/OCA2	rs12913832	T ^b	Brown	T	0.21	0.59	0.21	0.63	5.68E-39	0.29
OCA2	rs1800407	T	Blue	C	0.08	0.10	0.99	0.10	0.25	0.87
SLC24A4	rs12896399	$\mathbf{G}^{\mathbf{b}}$	Brown	G	0.43	0.63	0.012	0.66	1.02E-12	0.52
SLC45A2	rs16891982	C	Brown	C	0.02	_	0.94	0.16	5.62E-07	_
TYR	rs1393350	T	Blue	C	0.22	0.25	0.94	0.23	0.66	0.49
IRF4	rs12203592	Α	Blue	G	0.15	0.09	0.05	0.14	0.43	0.09

^a MAF, minor allele frequency (referenced to HapMap_CEU minor allele).

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Alleles with higher frequencies in the Italian and the Spanish population, becoming the major allele in both.

FpHWE p-value for Pearson's goodness of fit test for deviation from Hardy.—Weinberg equilibrium. p-value CEU and p-value TSI refer to the p-values of a Pearson's goodness of fit test comparing the Spanish minor allele frequency obtained from our samples with HapMap_CEU and HapMap_TSI frequencies respectively.

Bold in p-values denotes statistically significant results. Bold in minor alleles denotes alleles that become the major allele in our population.

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sample genotypes, being allocated different weights depending on whether an individual has two, one or no minor alleles. The probabilities were calculated according to the following formulas:

$$\pi_1 = \frac{exp(\alpha_1 + \sum \beta(\pi_1)_K x_K)}{1 + exp(\alpha_1 \sum \beta(\pi_1)_K x_K) + exp(\alpha_2 + \sum \beta(\pi_2)_K X_K)}$$

$$\pi_2 = \frac{exp(\alpha_2 + \sum \beta(\pi_2)_K x_K)}{1 + exp(\alpha_1 \sum \beta(\pi_1)_K x_K) + exp(\alpha_2 + \sum \beta(\pi_2)_K x_K)}$$

$$\pi_3 = 1 - \pi_1 - \pi_2$$

where x_k is the number of minor alleles of the kth SNP. The model's parameters, alpha and beta, were derived based on the genotypes of 3804 Dutch individuals in the model-building set of the previous study, and can be found in the same publication. These probabilities can be calculated using the macro provided in the supplementary material of Walsh et al. [13]. This prediction model classifies each individual as having brown, blue or an intermediate eye colour. Brown colour is considered when, given a specific IrisPlex SNP combination, the probability of being brown-eyed is higher than 0.5. Blue colour is predicted only when the probability of being blue-eyed is higher than 0.7. When, according to this six SNP genotype, the probability of being brown-eyed is less than 0.5 and, at the same time, the probability of being blue-eyed is lower than 0.7, the colour is classified as intermediate. The prediction relevance rank by SNP, in descending order of importance, was the following: rs12912832 first (most important), rs1800407 second. 12896399 third, rs16891982 fourth, rs1393350 fifth and finally rs12203592 sixth. Those individuals in which at least one SNP failed to be genotyped were excluded from this part of the study, since it was not possible to estimate eye colour probabilities, therefore a total of 493 individuals remained for subsequent analyses.

We assessed the accuracy of the prediction model by comparing the predicted eye colour with the real one, and then all samples were classified, for further analyses, as correctly predicted, darker than predicted, or lighter than predicted.

2.5. Statistical analysis

SPSS v19 was used to carry out the analyses. Bonferroni's correction was implemented as the method of adjustment for multiple comparisons. All *p*-values were two-sided and those lower than 0.008 were considered statistically significant.

For all polymorphisms studied, Fisher's exact test was used both to test for deviations from Hardy—Weinberg equilibrium (HWE) among the population sample and to compare differences in the minor allele frequency (MAF) distributions between cases and controls. HWE was rejected when the *p*-value was lower than 0.008 after Bonferroni's correction. Our data was compared with raw data obtained from HapMap_CEU and HapMap_TSI using unconditional logistic regression, in order to evaluate minor allele frequency differences between of our Spanish population and those available in HapMap.

Several analyses were performed to assess associations between genotypes and eye colour, and to evaluate the weight of each SNP, simultaneously allowing for various confounding factors. Genotype-related odds ratios (ORs) using the co-dominant model, their corresponding 95% confidence intervals (CIs) and associated *p*-values were estimated via unconditional logistic regression. This was done for each SNP as well as for gender. We also obtained results adjusted by possible confounding effects between SNPs or with phenotypic factors like gender after

including them all in a multivariate unconditional logistic regression analysis.

A meta-analysis was performed to compare our results to previously published data and check the robustness of our results. We searched for Caucasian populations in the literature containing data on both eye colour (classified in three categories that could be grouped in blue/non-blue eyes) and gender [10,17,18]. We conducted the meta-analysis on populations from Iceland, Holland, Australia and Poland along with the Spanish population analysed in this study, by applying a Cochran–Mantel–Haenszel test to the analyses performed on the total number of samples from all different populations, as previously described [19,20]. We set being male as a suspected risk factor for having blue eyes.

To evaluate differences in prediction accuracy between genders, we performed a 2-by-2 contingency table and calculated Chi-square values, *p*-values and associated odds ratios. We first considered the genotype of the *HERC2/OCA2* SNP and then we added the remaining IrisPlex SNPs and prediction algorithm. All samples were assembled in four different groups depending on gender and eye colour prediction: males correctly predicted, males wrongly predicted, females correctly predicted and females wrongly predicted. A sub-classification within the wrongly predicted groups was established based on whether the predicted eye colour was darker or lighter than the real colour. The groups obtained are described in Table 4.

3. Results 218

3.1. SNP allele distributions

Minor allele frequencies for each SNP in three different populations (HapMap_CEU and TSI, and Spanish) and *p*-values for the comparison between them and the Spanish population are detailed in Table 1, along with *p*-values of test of departure from Hardy_Weinberg equilibrium among our population. All polymorphism complied with Hardy_Weinberg equilibrium except SNP rs12896399 located within the *SLC24A4* gene.

We first observed that in both rs12913832 and rs12896399, the minor allele in the HapMap_CEU population becomes the major allele in both the Spanish and the HapMap_TSI populations. We were not able to test the difference between the Spanish and the Italian populations for SNP rs16891892, since this data is not available in HapMap.

Minor allele differences in the Spanish population differ significantly from HapMap_CEU frequencies in three SNPs: rs12913832 (*HERC2/OCA2*), rs12896399 (*SLC24A4*) and rs16891982 (*SLC45A2*). We observed that frequencies in our population were more similar to the ones in the HapMap_TSI population (Table 1).

3.2. Phenotype and genotype distribution

We classified all participants into three eye colour groups: brown, blue/grey and intermediate (intermediate phenotypes ranging from light green to light brown/hazel with shades of green). The Spanish population predominantly possesses brown eyes (75.47%), while blue-eyed people comprise 11.32% of the population, and there is a relatively high percentage (13.21%) of people with intermediate phenotypes (Fig. 1). When eye colour percentages are separated according to gender, percentages turn out to be different between males and females (Table 2). The percentage of blue-eyed females (8.5%) was notably lower than the one of blue-eyed males (14.71%) in the Spanish population, whereas browneyed females (78.45%) were considerably more common than males (71.43%). These differences presented statistically significant results (OR 1.34, 95% CI: 1.04–1.73, p-value 0.024) (Table 2).

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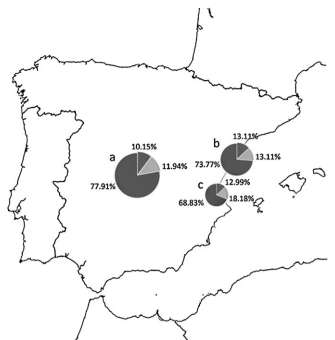


Fig. 1. Eye colour phenotype frequencies across three Spanish populations. Geographic locations: (a) Madrid (N = 332), (b) Castellon (N = 121), (c) Valencia (N = 82). Pie charts show the percentages of individuals with each phenotype. The size of each pie chart is proportional to the sample size. Stripped grey colour represents the frequency of blue-eyed individuals, dark grey represents browneyed individuals, and light grey represents the frequency of intermediate phenotypes.

In addition, we have performed a test to evaluate whether any of the other 5 IrisPlex SNPs are associated with gender, which would explain the differences in eye colour distribution between males and females. The results obtained demonstrated that none of the additional SNPs considered were associated with gender (data not shown).

We have performed a similar analysis using previously published eye colour frequency data from other Caucasian populations (Iceland, Holland, Australia, Poland) in which gender numbers were included [10,17,18]. The results of the present meta-analysis are shown in Fig. 2. When all individuals comprised in these studies, including the present one (a total of 10,292 individuals of Caucasian origin), are analysed together, the difference is extremely significant (p-value 1.56×10^{-10}). In addition, due to the fact that the Australian study by Duffy et al. [18] is actually derived from family data and not from unrelated individuals, which may have significantly distorted the results, we

performed an alternative analysis excluding the Australian data from the analysis (including only Iceland, Holland, Poland and Spain). The p-value achieved was 8.96×10^{-19} . Caucasian males definitely tend to have higher frequencies of blue/grey eye colour when compared to females who, in turn, display darker eyes (green or brown) much more commonly (see Fig. 2).

3.3. IrisPlex SNP association with blue eye phenotype

We performed a statistical analysis via unconditional logistic regression to assess the association between all six SNPs and blue eye colour. We tested the differences in genotype distribution between blue-eyed individuals and non-blue-eyed ones among 493 Spanish individuals. Since the eye colour differences between males and females described above suggested that gender could be a confounding factor, we decided to include gender in the analysis. As expected, HERC2/OCA2 SNP rs12913832 was found to be the major factor responsible for blue eye colour, with an OR of 0.03 (95% CI: 0.02–0.07, p-value 4.91 \times 10⁻²⁴). We also found a trend to significance for blue eye colour association with gender (OR 1.69, 95% CI: 0.96–2.97, p-value 0.068), meaning that male gender might be a factor contributing to blue eye colour (Table 3).

To assess possible confounding effects, a multivariate logistic regression analysis including SNPs and gender altogether was carried out. As expected, a strong association was found between HERC2/OCA2 and blue-eye colour $(p\text{-value }2.37\times10^{-18})$. SNP rs16891982 on SLC45A2 $(p\text{-value }2.5\times10^{-2})$ and, surprisingly, gender $(p\text{-value }1.2\times10^{-2})$ resulted in statistically significant associations with eye colour after adjustment for genotypes. Even though only HERC2/OCA2 association reached the Bonferroni threshold of 0.008, gender also showed a p-value nearing significance after correction. We also found marginal significant association with rs12203592, located on IRF4 (p-value 0.071) and with rs1800407 on OCA2 (p-value 0.075) (Table 3).

3.4. Accuracy of the prediction model

The IrisPlex predictive model was applied as described in Walsh et al. [13], and the prediction colour of each individual based on their genotypic profile of six SNPs located on six genes at different chromosome location was obtained. This colour prediction was compared with the real colour in each individual and the percentage of concordance was computed for all 493 individuals complying with the six-SNP IrisPlex genotype. Eye colour prediction count, percentages and concordance by phenotype and gender are detailed in Table 4.

The overall prediction accuracy is over 80% (83.38%) in the Spanish population. However, this prediction accuracy greatly varies depending on both eye colour and gender, predicting much

Table 2 Q6 HERC2/OCA2 genotypes and eye colour phenotypes among sampled Spanish individuals.

	Genotype $N=493^{c}$ (%)	Sex		<i>p</i> - <mark>V</mark> alue	OR (95% CI)	
		Female (n = 271) Male (n = 221)				
HERC2/OCA2 Genotype	CC 65 (13.19) CT 232 (47.06) TT 196 (39.76)	37 (13.60) 134 (49.26) 101 (37.13)	28 (12.67) 98 (44.34) 95 (42.99)	0.269	1.16 (0.89–1.51)	
		Sex				
	Phenotype $N = 535^{\circ}$ (%)	Female (n=297)	Male (n=238)	p- <mark>V</mark> alue	OR (95% CI)	
Eye colour phenotype	Blue/grey 60 (11.32) Intermediate 72 (13.46) Brown/black 403 (75.33)	25 (8.5) 39 (13.13) 233 (78.45)	35 (14.71) 33 (13.87) 170 (71.43)	0.024	1.34 (1.04–1.73)	

Q7 a p-Values and OR for t-student via unconditional logistic regression. OR (95% CI): odds ratio and 95% confidence interval.

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Percentages are shown in parentheses and calculated using as a total the N of each subgroup (total, female and male).

^c Note that the genotype and phenotype totals are different due to missing genotypes in some individuals.

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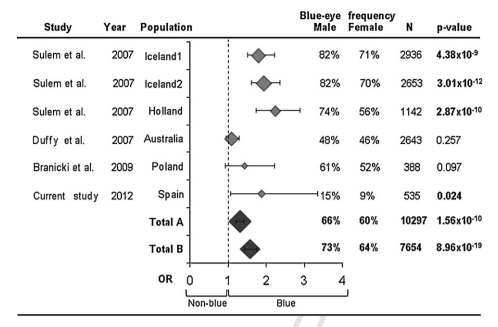


Fig. 2. Meta-analysis of the blue-eye colour trait in different Caucasian populations. N refers to the total number of samples analysed in each study. Total A and Total B p-values have been calculated using the Cochran,—Mantel—Haenszel test. OR: Odds ratio. Diamond shapes represent the overall odds ratio. The size of the diamond is proportional to the number of individuals, and error bars represent 95% confidence intervals. Light grey-filled diamonds represent data from individual studies, dark-filled diamonds represent the sum of all studies. Bold on p-values denote statistically significant results. Total A: Results achieved by taking into account all populations, including Australia. Total B: Results achieved by excluding Australian data from the analysis (including only Iceland 1 and 2, Holland, Poland and Spain). This alternative analysis was performed due to the fact that the Australian study of Duffy et al. [18] is actually derived from family data and not from unrelated individuals, which may have significantly distorted the results.

better brown-eyed individuals, 95.66% correctly predicted (with 95.60% for males and 95.71% for females) than blue-eyed individuals (76.36%). Overall, the accuracy for intermediate phenotypes is low, 23.18% (32.26% in males and only 15.79% in females). Although the ability to predict blue-eyed phenotypes

was considerable (76.36%), there was a remarkable male/female divergence, with 83.33% of blue-eyed females correctly predicted versus only 70.96% of blue-eyed males.

To further evaluate these eye colour mispredictions, we focused our attention on the SNP most associated to blue vs. non-blue

Table 3Genotypic association with blue eye colour.

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Gene	SNP	Allele/factor	Non-adjusted		Adjusted		
			OR (95% CI)	p-Value	OR (95% CI)	<i>p</i> -Value	
HERC2/OCA2	rs12913832	T	0.03 (0.02-0.07)	4.91 × 10 ⁻²⁴	0.02 (0.01-0.05)	2.37×10^{-18}	
OCA2	rs1800407	T	0.56 (0.25-1.25)	0.159	2.77 (0.90-8.35)	0.075	
SLC24A4	rs12896399	G	0.96 (0.65-1.42)	0.847	0.89 (0.49-1.59)	0.687	
SLC45A2	rs16891982	С	0.62 (0.33-1.15)	0.129	0.36 (0.15-0.89)	0.025	
TYR	rs1393350	T	0.90 (0.55-1.46)	0.669	1.35 (0.65-2.79)	0.416	
IRF4	rs12203592	A	1.38 (0.79–2.41)	0.260	2.12 (0.94–4.78)	0.071	
	Gender	Male	1.69 (0.96–2.97)	0.068	2.87 (1.26–6.54)	0.012	

^a p-Values for Fisher's exact test. OR (95% CI): odds ratio and 95% confidence interval.

 8 $^{\circ}$ Bold marks statistically significant results.

Table 4Eye colour phenotypes in the Spanish population, separated by sex and by predicted eye colour concordance (*N*=493).

Gender	Phenotype	Predicted eye colour			Total	Prediction-phenotype concordance	
		Blue	Intermediate	Brown			
Male (N=221)	Blue	22 (70.96)	2 (6.45)	7 (22.58)	31	22 (70.96)	
	Intermediate	3 (9.68)	10 (32.26)	18 (58.06)	31	10 (32.26)	
	Brown	1 (0.63)	6 (3.77)	152 (95.60)	159	152 (95.60)	
	Total	27	18	177		184 (83.25)	
Female (N=272)	Blue	20 (83.33%)	3 (12.50)	1 (0.16)	24	20 (83.33)	
	Intermediate	13 (34.21)	6 (15.79)	19 (50.00)	38	6 (15.79)	
	Brown	1 (0.48)	8 (3.81)	201 (95.71)	210	201 (95.71)	
	Total	34	17	221		227 (83.45)	

 Q^9 a Predicted using the mathematical model suggested by Liu, only in samples complying with all six IrisPlex SNP genotypes.

Please cite this article in press as: C. Martinez-Cadenas, et al., Gender is a major factor explaining discrepancies in eye colour prediction based on *HERC2/OCA2* genotype and the IrisPlex model, Forensic Sci. Int. Genet. (2013), http://dx.doi.org/10.1016/j.fsigen.2013.03.007

b Adjusted for all potential risk factors by including them on a multivariate logistic regression analysis, considering all six SNPs and gender as risk factors.

^dFull six-marker genotypes from 493 individuals were available for this analysis.

Percentages are shown in parentheses.

Prediction concordance percentages were obtained from all individuals within the considered phenotype.

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phenotype: HERC2/OCA2 rs12913832. Among all individuals homozygous for the HERC2/OCA2 minor allele (CC, associated with blue eyes), 92.30% were predicted blue by the IrisPlex algorithm. Of those, 30% were wrongly predicted blue, since they had either brown or an intermediate eye colour phenotype. Most importantly, 77.78% of the HERC2/OCA2 homozygotes wrongly predicted to have blue eves were females, a remarkable difference. The remaining 7.70% of CC individuals were correctly predicted as intermediate due to the genotypes of the five additional SNPs having alleles associated with darker phenotypes, excluding one female who actually had brown eyes. When considering only the HERC2/OCA2 heterozygotes, all were predicted as intermediate or brown-eyed by the IrisPlex algorithm, due to the considerable weight that the HERC2/OCA2 genotype has on eye colour. However, 5.2% of these heterozygote individuals were in fact blue-eyed, 67% of them actually being male - another gender-related difference. Moreover, a single HERC2/OCA2 TT homozygote individual, also a male, was detected who had actually blue eyes instead of brown eyes. None of the other five SNPs could explain this effect. We therefore observe a significant trend pointing to darker eye phenotypes in females and lighter in males when considering the HERC2/OCA2 genotype alone. These gender-related comparisons probably resulted in nonstatistical significant results due to the reduced size of the groups considered. Studies with larger sample sizes or populations with higher blue-eye frequencies would be necessary to analyse this effect in more detail.

4. Discussion

In this study, we have determined the genotypes of six pigmentation-related SNPs in 535 Spanish individuals in order to further our understanding of eye colour prediction based on genotypes in the Spanish population.

First, this work confirms the significant differences detected in allelic frequencies in pigmentation related genes according to latitude among European populations [10,14,16,21]. Three of the genotyped SNPs, rs12913832 (HERC2/OCA2 gene), rs12896399 (SLC24A4 gene), rs16891982 (SLC45A2 gene), had roughly reverse allele frequencies in Spain when compared to Northern European populations, more in the line of darker Mediterranean populations such as the Italians, Greeks and French [16,21]. This evident North-South variation in allele frequencies is indeed consistent with the correlation between solar UV radiation levels and skin pigmentation in Europe [22,23].

Although all six IrisPlex SNPs have been significantly associated with eye colour in populations of Northern European origin [4,8,10,13,14,16], in our study of a dark Mediterranean population we could only strongly associate with eye colour variation HERC2/ OCA2's rs12913832, and moderately IRF4's rs12203592 and the two coding SNPs in both OCA2 and SLC45A2, perhaps due to the relatively low frequencies of blue-eyed individuals in Southern Europe [16,24,25] (see Section 3 and Table 1).

However, the major finding of this work is indeed the seemingly important role played by gender in the association between genotypes (especially of HERC2/OCA2) and eye colour phenotype. This relationship is supported by the statistical significant results of the logistic regression analysis (Table 3).

The association between genotype, eye colour and gender is made particularly obvious when we consider the genotype of rs12913832 (in the HERC2/OCA2 region), the SNP more strongly correlated with eye colour. Previous studies have determined that rs12913832 accounts for most eye colour variation seen in Caucasians [5–7], and that the rest of the SNPs apparently implicated so far in eye colour, rs1800407 (OCA2 gene), rs12896399 (SLC24A4 gene), rs16891982 (SLC45A2 gene), rs1393350 (TYR gene), rs12203592 (IRF4 gene), rs1408799 in

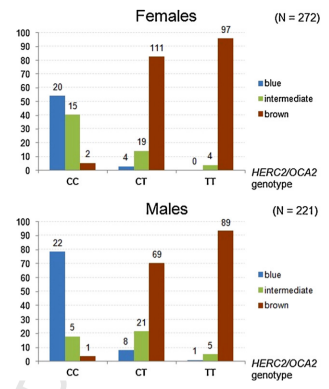


Fig. 3. Distribution/count of eye colour phenotypes according to their HERC2/OCA2 genotype and separated by gender, on a total of 493 Spanish controls. Percentages are calculated taking into account the total number of individuals within each HERC2/OCA2 genotype. The percentage of each phenotype (blue, intermediate and brown) is represented by bars of the corresponding colour. The number on the top of each bar represents the count of individuals in each category. For interpretation of the references to colour in this figure legend, the reader is referred to the web version of the article.)

TYRP, etc., have a relatively minor role [10,11,13,14,16]. But when, in our Spanish population, we focus solely on the genotype of rs12913832, we find that discrepancies between this SNP's genotype and eye colour phenotype are better explained by gender than by the addition of further SNPs. For example, most of the individuals (74%) in disagreement with a supposedly 'blue/grey genotype' (rs12913832 CC homozygotes that turn out not to have blue/grey eyes, and end up having a darker eye colour) are actually female. On the other hand, 70% of the individuals with blue/grey eyes not bearing the rs12913832 CC genotype (that is, having a lighter eye colour than suggested by their genotype), turn out to be male (see Fig. 3).

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Conversely, only 54% of the females with the rs12913832 CC genotype actually had blue or grey eyes, while most males with the CC genotype ended up having blue/grey eyes (79%). Even a relatively large fraction of the males carrying the CT or TT genotypes (4.7%) ended up having blue/grey eyes. On the other hand, almost all females (98.3%) carrying the supposedly 'darkeyed genotype' (rs12913832 CT heterozygotes and TT homozygotes) had indeed dark eyes (intermediate and brown) (All these data are shown in Fig. 3). This circumstance might also be expressed in terms of the sensitivity and specificity of a model. For example, we could actually state that, for the detection of blue eyes in males, the IrisPlex model demonstrates both a high specificity (most males predicted blue by the model actually had blue eyes), and a low sensitivity (a considerable proportion of males with blue eyes were not predicted blue by the model).

The conclusion or bottom line to be drawn from these results is that, given a particular HERC2/OCA2 genotype, males are more prone to have lighter eye colours than predicted by their

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genotypes, while females tend to have darker eye colours than predicted.

This effect is what may explain the fact that there seem to be comparatively higher frequencies of blue-eyed males than blueeyed females in populations of European origin such as Iceland [10], Holland [10], Australia [18] or Poland [17], as well as in this study (see Fig. 2). This appears to confirm the association between females and slightly darker eye colours and, conversely, males and lighter eve colours. This difference has been revealed to be highly significant (p-value 1.56×10^{-10} , see Results section), as demonstrated by the meta-analysis of available data from five different populations of Caucasian origin — Iceland, Holland, Australia, Poland and Spain (the p-value reaching 8.96×10^{-19} when the family-drawn biased Australian data was left out of the analysis). The results of this analysis (Fig. 2) show that males clearly and significantly exhibit blue/grey eyes more commonly than females, while green or brown eyes are indeed more frequent in females than males. Supporting this observation is the fact that in populations with very high blueeye frequency, such as Iceland or Holland, females show greater proportion of green eyes at the expense of blue eyes [10]. This green-eye phenotype could actually indicate a darkening of the blue eye colour genotype in females. However, in populations with predominantly dark eye colours such as the Spanish one, the effect of female eye darkening might simply be hidden by the common dark eye colour, being added or incorporated within the brown colour category, and thus showing no increment in female green-eve colour frequency.

These results apparently disagree with some previous studies suggesting that females around the world show on average lighter skin colour than males [22,26] based on a sexual selection hypothesis. In our opinion, our results do not challenge the evidence of lighter female skin, since both eye colour and specifically the HERC/OCA2 gene region do not seem to be associated with skin colour [10,27]. Furthermore, this male-female skin colour divergence does not necessarily have to be caused by genetic factors, and could be explained by physiological reasons such as male's considerable thicker skin and greater blood vessel content [28,29] and/or socio-cultural reasons (e.g. males spending more time outdoors and with less clothing than females). In addition, a recent GWAS by Candille et al. has pointed out that Caucasian males are actually more lightly pigmented in the unexposed skin than females [27].

Regarding the IrisPlex prediction algorithm, the model is much better at predicting blue eyes in females than males, due to the already mentioned excess of blue-eyed males not bearing the HERC2/OCA2 CC genotype. Although the model is equally good at predicting brown colour in males and females, it is true that it is worse at predicting intermediate/green colour in females than males, probably because more females bearing the 'blue eye genotype' (HERC2/OCA2 CC homozygous genotype) end up having green or intermediate eyes.

As a whole, the IrisPlex algorithm is quite good predicting brown eye colours, and relatively good at predicting blue/grey colours. But we have to take into account that there is a middle fraction, the intermediate/green coloured individuals, where the model shows great prediction difficulties. Moreover, this intermediate fraction seems to be slightly higher in the Spanish and other Mediterranean populations than in Northern European populations [16].

5. Conclusions

This study suggests that there exists a significant male-female eye colour divergence in the Spanish population as well as in other Caucasian populations. Additional studies in other populations of Caucasian origin are necessary to confirm this finding, mainly analysing the correlation between rs12913832 genotype and eye colour in relation to gender. This significant gender difference could suggest that there is an as yet unidentified gender-related factor contributing to human eye colour variation.

With regard to the IrisPlex prediction algorithm, the introduction of a 'gender factor' in the model that could take into account this gender divergence would allow for the prediction of slightly lighter eve colours for males as well as slightly darker eve colours for females. Probably, this gender factor would significantly improve the prediction success rate of the IrisPlex model or other models used to predict eye colour phenotypes from DNA genotypes.

Conflict of interests

Authors have declared that no conflicts of interests exist.

Acknowledgements

This study has been supported by grant number PI10_0405 from the Spanish National Health Ministry (Institute Carlos III). MPC is funded by the Education Council of the Generalitat Valenciana, VALi+d program (ACIF/2011/207). GR is funded by the Spanish Ministry of Health (CP08_00069). We would like to thank all the volunteers for giving their consent to participate in this study. We also thank the medical staff of Gregorio Marañón, La Paz and Ramón y Cajal Hospitals for collecting the samples. Kaspar genotyping was performed at the Central Medical Research Unit of the University of Valencia's School of Medicine.

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