Victoria University
School of Biomedical and Health Sciences

Blonde Hair Colour:
Classification, Characterisation and Genetic Associations
for use in Forensic Science

By
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A thesis submitted in fulfilment of the requirements for the
Degree of Doctor of Philosophy.

April 2010
Doctor of Philosophy Declaration

“I, Michelle Vaughn, declare that the PhD thesis entitled *Blonde Hair Colour: Classification, Characterisation and Genetic Associations for use in Forensic Science* is no more than 100,000 words in length including quotes and exclusive of tables, figures, appendices, bibliography, references and footnotes. This thesis contains no material that has been submitted previously, in whole or in part, for the award of any other academic degree or diploma. Except where otherwise indicated, this thesis is my own work”.

Signature: [Signature]

Date: March 24, 2010
This work is dedicated to my Grandparents

Charles and Robina Vaughn

&

Samuel James and Cecile Zacharias
AKNOWLEDGEMENTS

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ABSTRACT

The vast variety of normal pigmentation variation is a huge genetic puzzle and hair colour is one part of this puzzle. It is hypothesised that there are genetic variants specifically associated with an objectively defined Blonde phenotype. The first objective of this study was to examine the natural, adult hair colour variation among a population of European ancestry and to evaluate methods for both the measurement and classification of hair colours. The second objective was to characterise the microscopic and chemical features of hair colour in this sample and to determine how they relate to the macroscopic colour. The final objective was to compare genetic results with objectively measured and defined hair colour phenotypes in the search for genetic associations with ‘blonde’ hair colour. Having this information will assist in developing forensic predictive tests for physical features.

The presented data show that self-reported hair colours and observer-reported colours are similar; however, these categories are not necessarily the best way to categorize hair colour for quantitative research. Using a two-step cluster analysis, hair colour can be divided into categories or clusters based on spectrophotometric measurements in the CIE L*a*b* colour space and these clusters can be well discriminated from each other.

Being able to measure hair colour using digital image analysis may also be of interest. Using the CIE L*a*b* colour space, the hair colour of 196 individuals of European ancestry was measured by both reflective spectrophotometry and by digital image analysis (in V++). Only moderate correlations were found and it is concluded that, although more convenient, hair colour measurement from digital images has limited use in genetics or forensic science when accurate and consistent measurements are required, but may be useful in other less stringent fields such as intelligence or cosmetics.

Microscopic images of hair from 36 individuals were evaluated for colour (using the CIE L*a*b* colour space), hair thickness and the pattern of pigmentation. These measurements were compared to each other and to reflective spectrophotometric (RS) measurements of macroscopic hair colour. Despite a large ΔE* value, there is a strong relationship between macroscopic and microscopic L* values that would be useful if one was trying to predict the colour of a head of hair (macroscopic) from a strand (microscopic).
Eumelanin is a black/brown biopolymer pigment that is a major contributor to hair colour, along with other pigments. The eumelanin pigment component in hair was evaluated by quantifying the degradation product pyrrole-2,3,5-tricarboxylic acid (PTCA) by HPLC-UV. The amount of PTCA in the hair of 58 individuals was compared to macroscopic colour measurements using reflective spectrophotometric (RS) and microscopic colour measurements if available. There appears to be a significant relationship between several of the macroscopic and microscopic colour components and the amount of PTCA quantified, such as with the macroscopic RS $b^*$ value and with the microscopic L*.

Tests for genetic associations were carried out on both categorical variables (clusters) and quantitative variables ($L^*$, $a^*$ and $b^*$ colour values). In the categorical analysis, five of 58 SNPs show various significant associations with Fair/Blonde hair. These SNPs are in the genes MC1R (R151C), DTNBP1 (H297Y), TYR (R402Q), ATP7B (R832K) and MYO7A (C1628S). MC1R (R151C) and TYR (R402Q) were found to be associated with Red hair. Associations were also found with Dark hair in this sample; a SNP in the gene MATP (L374F) was significantly associated with dark hair in both the allelic and dominant model. Two SNPs in ATP7B (S406A and V456L) were also found to be associated in a dominant model with Dark hair.

In the quantitative variable analysis, polymorphisms in the genes TYR, MATP, DTNBP1, ATRN, MC1R, MLPH, MYO7A and MYO5A were found to be associated with $L^*$ values and these results tend to correspond with genes that were associated with light or dark hair in the categorical analysis.

Future directions of this research could include similar studies with increased sample sizes, greater investigation into the cell biology of hair and the genotyping of more SNPs.

In conclusion, this study successfully met its aims of objectively classifying and characterising blonde (or fair) hair colour and finding genetic associations with blonde hair colour. The results obtained in this study will benefit future research into the genetics of hair colour for use in forensic science and possibly in other areas of pigmentation and forensic research.
PUBLICATIONS

Peer-Reviewed Articles:


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Hair colour measurement by reflective spectrophotometry and digital image analysis

Michelle Vaughn, Elizabeth Brooks, Bruce Comber, Roland van Oorschot and Swati Baindur-Hudson

Characterisation of microscopic hair colour and texture and a comparison with macroscopic colour

Michelle Vaughn, Domenico Caridi, Roland van Oorschot and Swati Baindur-Hudson

Eumelanin characterisation in hair compared to colour measurements

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Hair colour measurement and classification
ABBREVIATIONS

2-S-CD  2-S-Cysteynldopa
3-AHP  3-amino-4-hydroxyphenylalanine
3-AT   3-aminotyrosine
DHI    5,6-dihydroxyindole
DHICA  5,6-dihydroxyindole-2-carboxylic acid
5-S-CD 5-S-Cysteynldopa
AIM(s) ancestry informative marker(s)
BGA biogeographical ancestry
ΔE* Change in Colour
ΔL* Change in Lightness
CCD charge coupled device
CODIS Combined DNA Index System
CIE Commission Internationale de l'Eclairage
Co cortex
Cu cuticle
CYM Cyan-Yellow-Magenta
DNA deoxyribonucleic acid
ddH2O distilled and deionised water
DI digital image
E# Euclidian Cluster Number
FBI Federal Bureau of Investigation
FD fluorescence detection
g gravities
HWE Hardy-Weinberg Equilibrium
HPLC high performance liquid chromatography
HSV Hue, Saturation, Value
LD linkage disequilibrium
LC/MS liquid chromatography/mass spectrometry
LL# Log-likelihood Cluster Number
mAU milli Active Unit
Md medulla
<table>
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<td>MSHR</td>
<td>melanocyte stimulating hormone receptor</td>
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<tr>
<td>MF</td>
<td>microfibril</td>
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<td>µL</td>
<td>microlitre</td>
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<tr>
<td>mL</td>
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<tr>
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<tr>
<td>NCIDD</td>
<td>National Criminal Investigation DNA Databank</td>
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<tr>
<td>nm</td>
<td>nanometre</td>
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<tr>
<td>ORC</td>
<td>observer-reported colour</td>
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<tr>
<td>OR</td>
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<tr>
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<tr>
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<td>parts per million</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
</tr>
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<td>PTCA</td>
<td>pyrolle-2,3,5-tricarboxylic acid</td>
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<tr>
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<tr>
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</tr>
<tr>
<td>RGB</td>
<td>Red-Green-Blue</td>
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<td>reflective spectrophotometry</td>
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Chapter 1     Introduction

1.1     General Introduction

1.1.1     Introduction

“Color is so much a matter of direct and immediate perception that any discussion of theory needs to be accompanied by experiments with the colors themselves.”

-- Walter Sargent (1868-1927)[1]

Pigmentation in humans has been observed and studied for at least 4000 years [2] and is an integral part of our physical health and cultural identity. Scientists have been interested in the inheritance of pigmentation traits for over a century and have been studying the genetics of pigmentation for more than 50 years [3]. Efforts in the past have concentrated a great deal on pigmentary disorders [4, 5] and on murine models of pigmentation, where over 150 alleles at more than 90 genetic loci have been discovered so far [6].

Study of the normal pigmentation variation in humans is more recent and has been investigated by medical scientists [7], anthropologists [8] and, more recently, by forensic science researchers for the prediction of visible phenotypes to be used as an investigative tool [9]. In the field of forensic science, iris colour has received a great deal of attention, as has the genetic determination of biogeographical ancestry (BGA) [10-12]. An American company, DNAPrint, has developed tests for BGA that have been used in casework in the United States [12] and are developing a kit for the prediction of iris colour [9, 13].

Hair colour has received much less attention in the study of pigmentation, with most work on hair colour being done as a consequence of, or in correlation with, another pigmentary trait (such as iris colour or melanoma risk phototype). Parts of this investigation have been done by researchers in the vastly different fields of medical science [7, 14, 15], the cosmetics industry [16, 17] and forensic science [18], as well as in chemistry and physics [19, 20]. The reported project is specifically concerned with genetic determinants of blonde hair colour.
Many parties may be interested in research on blonde hair colour. Blonde hair colour has been implicated as a melanoma risk factor [7] and genetic determinants may be interesting to medical, anthropological and cosmetic researchers. The prediction of physical traits for use in forensic investigations is a primary interest in this research and hair colour is one of many traits that may assist investigators in limiting a suspect pool. The UK Forensic Science Service has already implemented a test with 84% accuracy for predicting red hair [18].

To produce an accurate and valid test for physical characteristics, the genetic basis of these characteristics must first be solidly understood. The vast variety of normal pigmentation variation is a huge genetic puzzle and hair colour is one part of this puzzle. By sequentially examining smaller components, the larger picture of human pigmentation variation will emerge.

1.1.2 Research Outline and Objectives

It is hypothesised that there are genetic variants specifically associated with an objectively defined Blonde phenotype. This project aims to first characterise and define the phenotype under study (‘blonde hair’) and then to evaluate DNA sequence variants that may be associated with this phenotype. These variants may be useful in predicting the hair colour of an unknown subject.

A review of the scientific literature relevant to forensic science and the prediction of physical features can be found in Chapter 2. Further discussion of the background literature that is relevant to each experiment can be found in Chapters 3 through 7.

As well as incorporating a review of the applicable literature, Chapters 3 through 7 discuss the methods and results of each experiment. Chapters 3 and 4 are concerned with objective measurement and definition of hair colour phenotypes. Chapters 5 and 6 involve a more thorough characterisation of a subset of samples and Chapter 7 describes the genetic analysis.

In Chapter 3, natural hair colour is measured by reflective spectrophotometry and these objective measurements are compared to observer and self-reported colours and are
classified into defined groups by cluster analysis. Chapter 4 involves the comparison of the reflective spectrophotometry measurements with colour measurements obtained by digital image analysis.

**Objective 1** - Examine the natural, adult hair colour variation among a population of European ancestry and evaluate methods for both the measurement and classification of hair colours.

**Why?** To assist in the development of standardisation in hair colour research for this project, for future research and in forensic science.

Chapter 5 reports the findings of a microscopic hair examination of a subset of individuals and compares microscopic features of hair colour and hair texture with the macroscopic colour. A chemical analysis of the pigments in hair is described in Chapter 6 and the chemical composition of hair pigments is compared to macroscopic and microscopic colour measurements.

**Objective 2** – Characterise the microscopic and chemical features of hair colour in this sample and how they relate to the macroscopic colour.

**Why?** To increase the understanding of hair colour biology and chemistry; this will assist in both traditional forensic hair examination and in designing and interpreting genetic studies.

Chapter 7 examines the relationship between hair colour and genotype at the site of 58 single nucleotide polymorphisms. Associations within the sample are presented and additional findings will be discussed.

**Objective 3** – To compare genetic results with objectively measured and defined hair colour phenotypes in the search for genetic associations with ‘blonde’ hair colour.

**Why?** Having this information will assist in developing forensic predictive tests for physical features.
A final discussion of the research, its findings and possible future directions will be presented in Chapter 8.
Chapter 2   Background and Literature Review

2.1  Forensic Biology

2.1.1  Forensic Hair Examination

Hair is commonly encountered as physical evidence at crime scenes [21, 22]. Traditional forensic hair examination involves examining the specimen under a microscope and observing characteristics such as colour, length and diameter of the hair, as well as the distribution of pigment granules and other morphological features of the hair that may assist in determining the source of the hair [21-24].

An expert hair examiner may be able to determine a great deal of information about the contributor of a hair and will compare the features of a ‘questioned’ hair to those of a hair sample from a ‘known’ source. The examiner may be able to determine if the hair is human and if not, what species it belongs too [22]. Human hair will be evaluated for its length and shape, which will help determine the racial origin of the hair and the area of the body it is from. The tip of the hair (e.g. cut or abraded), the macroscopic colour of the hair and any evidence of disease will also be evaluated [22].

Microscopic features of the hair will also be examined. This will include evaluations of the hairs’ colour, diameter, cross-section, pigment features, medulla classification, cortical features and cuticle features [22]. An in-depth description of forensic hair examination can be found in James Robertson’s book ‘Forensic Examination of Human Hair’ (1999) [22].

Figure 2.1 gives an example of checklists that is used by the Australian Federal Police forensic hair examiners.
<table>
<thead>
<tr>
<th>HAIR NUMBER</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>MACROSCOPIC FEATURE</td>
<td>r</td>
<td>r</td>
<td>t</td>
<td>r</td>
<td>t</td>
</tr>
<tr>
<td>LENGTH</td>
<td>cm</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SHAFT PROFILE</td>
<td>Straight</td>
<td>Wavy</td>
<td>Curly</td>
<td>Peppercorn</td>
<td></td>
</tr>
<tr>
<td>COLOUR¹</td>
<td>Colourless</td>
<td>Yellow</td>
<td>Brown</td>
<td>Reddish</td>
<td>Black</td>
</tr>
<tr>
<td>ROOT</td>
<td>Absent</td>
<td>Club</td>
<td>Ribbon</td>
<td>Sheath Present</td>
<td></td>
</tr>
<tr>
<td>TIP</td>
<td>Natural Taper</td>
<td>Cut</td>
<td>Rounded</td>
<td>Frayed or Abraded</td>
<td>Split</td>
</tr>
</tbody>
</table>

General Description and Comments:

¹ A assessed with a stereo microscope using standardised illumination
B basic colour to be qualified by shade or depth of colour. Light (L), Mid (M) or Dark (D)
C note artificial colouring

Examined by: __________________________

Notes by: __________________________

Day: __________________________

Date: __________________________

Time: __________________________
### AUSTRALIAN FEDERAL POLICE

**FORENSIC SERVICES, SCIENTIFIC**

**EXAMINATION RECORD - Sheet 2**

<table>
<thead>
<tr>
<th>HAIR NUMBER</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>MICROSCOPIC FEATURE</td>
<td>r</td>
<td>r</td>
<td>r</td>
<td>r</td>
<td>r</td>
</tr>
<tr>
<td>SHAFT DIAMETER</td>
<td>max units at × 40</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PIGMENT DENSITY</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PIGMENT DISTRIBUTION</td>
<td>Uniform</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(across hair shaft)</td>
<td>Towards Medulla</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PIGMENT AGGREGATE SHAPE</td>
<td>Streaked</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SHAPE</td>
<td>Clumped Oval</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PIGMENT GRANULE SHAPE</td>
<td>Fine</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SIZE</td>
<td>Oval/Oblong</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PIGMENT AGGREGATE SIZE</td>
<td>Fine</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GRANULE SIZE</td>
<td>Medium</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OVOID BODIES²</td>
<td>Fine</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MEDULLA DISTRIBUTION</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MEDULLA TYPE</td>
<td>Opalescent</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CORTICAL FUSI²</td>
<td>Not Visible or Smooth</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CORTICAL TEXTURE</td>
<td>Visible or Coarse</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CUTICLE</td>
<td>Thickness units × 40</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CUTICLE OUTER</td>
<td>Smooth</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Colour</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cracked</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

² Where present their shape, size and distribution both along and across the hair shaft may have some value when used as comparative features.

Examined by: ________________________  Checked by: ________________________

Notes by: ________________________  Day: ________________________  Date: ________________________  Time: ________________________

---

**Figure 2.1** Hair examination checklists for the Australian Federal Police
The American National Research Council recently published a document entitled “Strengthening Forensic Science in the United States: A Path Forward” [25]. This document expresses the need for standardised terminology and reporting of results in forensic sciences, as well as for an improved scientific basis for forensic science techniques and knowledge [25]. In contrast to DNA evidence, which uses standardised and objective protocols to determine genotype data and then uses a statistical calculation for identification, traditional hair examination still involves an expert’s opinion and a hair cannot currently be conclusively attributed to an individual. The features that are highly variable between individuals are those which are difficult to capture easily or objectively, such as pigment pattern and colour. However, there are efforts being made to develop objective and validated methods for the forensic examination of hair and to increase the value of hair as evidence [24, 26]. One of these efforts involves the use of pigmentation patterns and colour measurements for identifying the contributor of a hair [27].
2.1.2 DNA

Deoxyribonucleic acid, or DNA, is a molecular instruction book that resides in almost every cell of our body. This molecule contains the information about what blood type a person is, their predisposition for many diseases and, theoretically, what one looks like. The DNA in each individual is unique, with the exception of identical twins. This uniqueness allows for a powerful means of determining the source of biological material found at crime scenes or the identification of victims of crime. DNA identity testing has become routine in the course of criminal investigations [28, 29]. By comparing to a sample of known origin, forensic biologists determine the probability that an individual was the source of a particular DNA sample; whether it is a blood stain, skin under the fingernails of a victim or the paternity of a child. The word “match” is used in conversation, however; it is not quite correct. A more correct phrase would be to conclude that a subject ‘can not be excluded as the contributor of the DNA’ and the likelihood of someone else being the contributor is expressed as a probability [28]. A full, good-quality, DNA profile produces probabilities such as one in trillions – a conclusive result.

This profile is typically arrived at by the testing of genetic markers called Short Tandem Repeats (STRs). Short Tandem Repeats involve regions of the genome that do not code for specific proteins (non-coding regions) and where short sequences of nucleotides are repeated. For example, the STR called TH01 is used in forensic science and consists of five to eleven repeats of the sequence AATG [21, 28]. As for any genetic marker, there are two copies; one on the maternal chromosome and one on the paternal chromosome, which may have the same or a different number of repeats. Therefore, in this example, an individual may have, for that marker, a TH01 profile of 6 and 8 repeats.

STRs are typically tested by capillary electrophoresis in the laboratory via strictly controlled and validated protocols where the size of the markers (and therefore, the number of repeats) is determined [28]. The number of markers tested will depend on the case involved and the exact markers tested will depend on the laboratory. A well known system is the Combined DNA Index System (CODIS) used by the FBI. This consists of 13 STR markers that are completely independent of one another and that have known population frequencies [28]. In addition to the STR marker, one coding
sequence is typically used in forensic DNA profiling to determine the sex of the individual. The amelogenin gene (which codes for an extracellular matrix protein in tooth enamel) is shorter by six base pairs on the X chromosome compared to the Y and so a female will show one fragment size, a male will show two [28]. Australian forensic scientists use a similar system for the National Criminal Investigation DNA Databank (NCIDD at www.crimtrac.gov.au).

It is important to note that in normal DNA profiling for forensic purposes, only non-coding genetic markers are used (except for amelogenin) and, besides sex, nothing is known about the individual, it is only a test for identity. In such profiling, samples from suspects, criminal databases and victims are compared to samples from the scene of a crime. If there is no comparison sample available (such as a suspect sample or database hit), the DNA sample is of limited investigative value.

A typical DNA profile of a female using nine STRs and amelogenin is shown in Figure 2.2.
Figure 2.2 A nine-locus (+ amelogenin) profile of a female that may be used for identification

2.1.3 Future Directions in Forensic Science

The field of forensic science is changing rapidly, with heightened efforts in scientific research as a response to increased demands from the courts and investigators for useful, validated techniques. Evidentiary forensic science involves the comparison of samples of ‘questioned’ origin to those of ‘known’ origin; whether it is a DNA sample, a hair or other trace evidence or a tool mark impression.
A newer area of forensic science is ‘forensic intelligence’ – forensic science that assists with an investigation; prior to an arrest or court-room activity [30]. Science may be able to provide the police with information that can direct them to a particular individual or location. Some examples of this in use today include psychological and geographic profiling as well as the linking of crime scenes through trends in physical evidence such as foot prints, tool marks and DNA [30].

Using genetic information to predict what someone looks like is one area of investigative forensic science that has gained a great deal of interest in recent years. This is alternatively called ‘phenotype profiling’, ‘physical feature prediction’ or ‘molecular photofitting’. Phenotype profiling is based on the idea that our physical features (such as skin, hair and eye colour, baldness patterns, height, etc.) are coded for in our DNA and if one was to test for these variations, a description of an individual could be produced and offered to investigators. Having this information would assist the police in narrowing a suspect pool or beginning a search for a perpetrator, a victim or even a witness [9, 31].

2.2 Phenotype Profiling

2.2.1 Ancestry

One of the more developed areas of research in phenotype profiling is the estimation of a person’s biogeographical ancestry (BGA) from their genetic sequence [9]. Ancestry Informative Markers (AIMs) are genetic markers (usually single nucleotide polymorphisms or SNPs) that vary in frequency between populations that were once geographically isolated from one another due to genetic drift [9]. By genotyping many of these markers a statistical estimate of a person’s genomic ancestry can be determined. The strength of the estimate may depend on how many markers are used in the calculation and how informative the particular markers are for ancestry [9, 32, 33].

It is important to differentiate between genealogical ancestry and genomic ancestry. Genealogical ancestry depends on the population one’s ancestors derived from (fixed by history) where genomic ancestry is the average level of admixture of an individual [9]. For example; a person may have 3 grandparents of European origin and 1 of African
origin. Therefore there genealogical ancestry would be 75% European and 25% African, but their genomic ancestry (or BGA) may not be the same. Due to independent assortment and recombination of chromosomes, this person may have more or less than or equal to 25% African genomic ancestry. Figure 2.3 illustrates a hypothetical example where an individual has 75% European and 25% African genealogical ancestry, but approximately 61% European and 39% African genomic ancestry due to random assortment of chromosomes in meiosis. This example does not take into account any recombination of chromosomes.

Figure 2.3 A hypothetical pedigree of one individual who has 75% European and 25% African genealogical ancestry, but approximately 61% European and 39% African genomic ancestry due to random assortment of chromosomes in meiosis. This example does not take into account any recombination of chromosomes.

Accurate measures of biogeographical ancestry are necessary to assist clinicians, biomedical researchers and forensic scientists [9]. There is often variation in drug responses across various populations and knowledge of a patient’s BGA will assist a clinician in determining the correct treatment [9]. Assessment of BGA also increases the accuracy of medical genetic studies by sorting out which variants are related to ancestry rather than disease [9] (this may also apply to the study of complex non-
disease traits such as pigmentation). Forensic DNA analysts are interested in measures of BGA to assist with likelihood calculations for identity testing and for inferring physical characteristics of an individual [9].

### 2.2.2 Indirect Phenotype Inference

Physical traits such as pigmentation and craniofacial morphology can be inferred from ancestry using an indirect method of phenotype inference [9]. In this case, ancestry, as calculated by testing AIMs, is used as a proxy for the net effect of phenotypically functional loci and can give information on traits that have a strong association with ancestry, such as pigmentation [9]. For example, an individual with 80% African ancestry is likely to have darker skin than an individual with 30% African ancestry, however, the range of informative admixture values must be determined empirically [8, 9]. Frudakis (2008) discusses the use of admixture databases to assist with the interpretation of indirect phenotype inference [9].

### 2.2.3 Direct Phenotype Inference

Direct phenotype inference involves the evaluation of genetic information that is directly related to the trait being examined [9]. This can be used if the genetic cause of the trait is understood and results in higher quality information than an indirect inference, however this information is much harder to come by [9].

Pigmentation may have a relatively more straightforward genetic mechanism than something like cardiovascular health due to more obvious selection pressures and this makes it more amenable to study the direct genetic causes of pigmentation traits [9], however there is still a great deal of information to be gathered and understood.

To develop a predictive test for a pigmentation feature, Frudakis (2008) presents four steps that may be considered. First, associations must be identified between genetic markers and phenotypic traits. Next, associations between certain genetic polymorphisms and certain trait values must be established. The inferential method for predicting an individual’s trait value must be then validated and finally, the trait can be inferred directly [9].
The presented research is primarily concerned with the first step in developing method for the direct inference of hair colour from genetic information (finding associations) and the literature in this field will be discussed next.

2.3 **Direct Inference of Hair Colour**

2.3.1 **Human Hair Colour**

Hair colour is one physical feature of interest to those studying the genetics of physical traits for phenotype profiling. In humans, hair colour is a continuous trait [9], but one that is usually broken into categories such as “blonde” or “brown” for reference. Hair colour in humans is the result of two main pigments, the black eumelanin and red pheomelanin, being transferred from the melanocytes where it is produced to the keratinocytes of the hair shaft. Details of the pigmentation process (as it is thought to occur in the epidermis) and the genes thought to be involved will be discussed in later sections of this thesis; however an overview of the process is illustrated in Figure 2.4.

The final colour of the hair depends on the amount and type of melanin, which are related to the shape, size and type of melanosomes (the organelles where melanogenesis occurs) and the activity of Tyrosinase (the primary enzyme responsible for melanin production) [9]. One can see from Figure 2.4 that there are many genes involved in the control of this process and there are likely more genes that have not yet been associated with pigmentation at all.
Figure 2.4 Graphical representation of a melanocyte showing some important proteins involved in melanocyte development and the pigmentation pathway (as thought to occur in the epidermis), including the following: 1) the tyrosinase enzyme complex (TYR, TRP1, DCT) located on the membrane of the melanosome, which is responsible for the enzymatic conversion of the amino acid tyrosine into melanin; 2) other proteins located within the melanosomes that play a critical role in melanogenesis (MATP, P gene, SILV, SLC24A5); 3) the signalling pathways affecting the regulation of melanin synthesis, including hormones and receptors (α-MSH, MC1R, ASIP, ATRN); 4) the transcription factors involved in melanin production (PAX3, MITF, SOX10); 5) proteins involved in melanosome transport (MYO5A, MYO7A, RAB27A) and melanosome construction/protein routing (CHS1, HPS1-6); and 6) developmentally important ligands (EDN3, KITLG) and receptors (KIT, EDNRB) which control melanoblast migration and differentiation [34].
Research into finding genetic associations with hair colour has been less successful and is a less developed field than research in the areas of skin and iris pigmentation. Iris pigmentation is thought to be primarily associated with variations the OCA2 gene (also known as the P gene) shown in Figure 2.4 [9, 10, 35]. Iris colour has been well characterised, phenotype data bases have been created and luckily, while several genes are likely involved in iris pigmentation the OCA2 gene seems to have the greatest effect [9].

There are several possible reasons that the search for genetic associations with hair colour has been less successful and Section 2.3.3 will discuss these issues. They are also summarised in Frudakis (2008) [9].

2.3.2 Genetic Studies of Hair Colour

The first association of a genetic variation with hair colour was in 1995 where variations within the gene MC1R, which codes for the melanocortin 1 receptor (the melanocyte-stimulating hormone receptor or MSHR), were found to be associated with red hair [36]. The interest in red hair and MC1R continues [14, 37-42], perhaps due to its associations with skin-cancer risk and perhaps due to the fact that it is reasonably simple to study (MC1R is a small gene with very clear effects). Due to an obvious phenotype and a simple genetic mechanism, testing for red hair has also become one of the first direct phenotype inferences used in forensic science [18].

Research into the genetics of non-red hair colours and genes other than MC1R is less well established. One polymorphism in the 3’ untranslated region of the human agouti signalling gene (ASIP g.8818A>G) has been associated with darker pigmentation in African populations [43] and with darker hair and eye colour in Europeans [44]. Two polymorphisms in the MATP gene – also known as SLC45A2 - (Phe374Leu and Glu272Lys) have been associated with darker pigmentation (including hair) in European populations [45].

Recently (December 2007), an Icelandic company called ‘deCODE genetics’ published a very large study on the genetics of various pigmentation traits, including hair colour [46] which found several associations with Red hair, as well as Blonde (vs. not-Blonde) in genes such as MC1R, OCA2, TYR, KITLG and SLC24A4.
2.3.3 **Issues in the Study of Human Hair Colour Genetics**

One major issue in the study of human hair pigmentation is that different genes may be involved than are involved in iris and skin colour [9]. Pigmentation has traditionally been studied in mice [47, 48] and in human epidermis [4]. It is now known that there are fundamental differences between mouse and human pigmentation and between skin and hair pigmentation. For example, pheomelanin, the red-yellow pigment that is found in high proportions of red-haired humans, sometimes appears yellow in mice and these mice (*recessive yellow*) have almost no eumelanin [47], which does not occur in humans (see also Box 6.1) [20, 47].

The differences between skin and hair pigmentation are also being elucidated. It is well known that a person can have dark hair and fair skin and sometimes even fair hair and dark skin (such as is found amongst some Melanesian and Australian aboriginal populations) [49]. This is possibly the result of the process of melanin degradation being different in hair and skin and may be due to other means of control [49]. Recent gene expression studies [50] have shown that at least one protein (Tyrosinase-related protein Type 2) is expressed in epidermal melanocytes, but not in hair follicle melanocytes.

In addition to different genetic mechanisms, hair colour may be affected by the environment (external and internal) more so than skin or iris pigmentation [9]. It is widely known that hair colour can change throughout a person’s life. Hair colour often changes in puberty and as a person ages. Hormonal regulation of melanogenesis is thought to be involved in these processes [51] and hormonal disorders or certain medications may have additional effects on hair colour. Hair may also change in response to external environmental conditions, such as sunlight. Photo-lightening of hair has a great deal of anecdotal evidence, but few scientific studies. Takahashi and Nakamura (2004 and 2005) measured the photo-lightening of blonde and red hairs, and found that strands of both lightened in response to visible and ultraviolet light, by a maximum of about five $\Delta L^*$ units [16, 17] ($\Delta L^*$ and the CIE $L^*a^*b^*$ space will be discussed in Chapter 3). Larger studies on heads of hair rather than strands may assist in quantifying the photo-lightening affect of sunlight on hair.
Hair may also be coloured artificially and a person’s memory may not be accurate when asked to self-report hair colour later on. Each of these factors may introduce error into genetic studies of hair colour and will present problems if predictive tests are put into use in the future; however these factors can be controlled for in studies at this point in the research process by choosing subjects who are healthy and have their natural, adult hair colour at the time of sampling.

Unknown genetic interactions may also complicate the study of hair colour. There is most likely epistatic interactions between genes involved in hair pigmentation and they may be different than any discovered in epidermal or iris pigmentation. For example, a mutation in the MATP gene (g.8818A>G) is thought to suppress red-hair causing mutations in MC1R and leave an individual with darker hair [44].

Another major obstacle in the genetic study of hair colour, as previously mentioned, is the difficulty in defining the hair colour phenotypes. Studies of hair colour often ask the subject to report their own hair colour or the observations of the researcher are used [38, 45, 46] and while objective measurements are becoming more common, they are sometimes weighted towards one hair colour [52] or involve children, whose hair colour may yet change [41]. Objective hair colour measurements are difficult to obtain on large samples; however standardisation in methods of hair colour description and classification will increase comparability of various research studies and will assist in communication between research groups, forensic scientists and investigators.

The relationship between genetic variations and hair colour may also be more closely related with different types of hair colour measurements [53], such as chemical composition or microscopic colour and understanding the relationships between these measurements and macroscopic hair colour will assist in defining the hair colour phenotypes.

As mentioned above, the deCODE genetics study found several associations with Red hair, as well as Blonde (vs. not-Blonde) in genes such as MC1R, OCA2, TYR, KITLG and SLC24A4. This study lends support to the idea that several, less penetrant alleles are responsible for non-red variation. The odds ratios (a measure of the effect a mutation has) for blonde hair are lower than what is typically found for associations with red hair. The authors (Sulem et al.) note that no objective measurements of hair
colour were carried out. This study included almost 7000 individuals and illustrates two problems in analysing hair colour genetics – that hair colour is likely controlled by several low-penetrance alleles (which are harder to find in a case-control study) and that objective measurements of hair colour (for consistency and accuracy) are difficult or impossible in such large studies. In the future, it may be possible to extrapolate information from the measurement of smaller data sets, such as that presented in this project, by constructing a visual database of hair colour groups.

2.3.4 Continuing the Study of Hair Colour

As outlined in Section 1.1.2, each of the following chapters measured hair colour at a different level or in a different way. Further literature, relevant to a specific experiment, will be included in each chapter. Macroscopic colour, microscopic colour, chemical composition and genetic variation will be examined, analysed and discussed and it is hoped that each section individually and when considered together will provide information on hair colour that will be interesting and useful to those in the fields of genetics and forensic science.
Chapter 3   Hair Colour Measurement and Variation\textsuperscript{1}

3.1   Introduction

3.1.1   Colour and Perception

Colour is defined by the Random House Unabridged Dictionary as “the quality of an object or substance with respect to light reflected by the object, usually determined visually by measurement of hue, saturation, and brightness of the reflected light”\textsuperscript{[54]}.

When light of a certain wavelength is reflected from an object (the dominant wavelength), it appears to the human observer as colour \textsuperscript{[55]}. Visible light or “white light” is made up of electromagnetic radiation wavelengths in the 380 nanometre to 780 nanometre range. Figure 3.1 illustrates how these colours relate to other forms of electromagnetic radiation and what colours result from the visible spectrum.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{em_spectrum.jpg}
\caption{Energy spectrum including the wavelengths of visible light that appear as colours\textsuperscript{2}}
\end{figure}

The human eye is an incredibly complex structure and is capable of detecting wavelengths of light in the visible spectrum \textsuperscript{[55]}. For the purposes of this research, the

\textsuperscript{1} An earlier version of this experiment was published in the American Journal of Physical Anthropology Vaughn, M. et al. 2008: 137(1) Pages 91-96. Please see Appendix 4
\textsuperscript{2} http://www.antonine-education.co.uk/physics_gcse/Unit_1/Topic_5/em_spectrum.jpg
colour sensitivity of the eye will be briefly discussed. In the retina (which covers most of the interior of the eye) there are photosensitive receptors called rods and cones that feed stimuli to the relevant neurons and nerves. Rods give monochromatic vision while cones contribute to colour vision [55], although all contribute to three ‘channels’ of vision; achromatic (brightness) and two chromatic channels (red-green and yellow-blue) [55, 56]. This is referred to as having a trichromatic system of colour vision.

3.1.2 Colour Measurement

To objectively measure hair colour, a method of measuring colour must first be chosen. As mentioned in the above definition, colour can be described in terms of its hue \(H\), saturation \(S\) and brightness value \(V\) [55] as shown in Figure 3.2. This type of model is referred to as a tri-stimulus colour model or colour space. Other examples include the additive Red-Green-Blue (RGB) colour model that is often used to display images on a computer screen (Figure 3.3) and the subtractive Cyan-Magenta-Yellow (CYM) model (Figure 3.4) that is found in some printers [57]. RGB and CMY colour models are useful, however are difficult to associate with human perception of colour and are mainly used for devices.

![Figure 3.2 The HSV colour space showing hue (H), saturation (S) and brightness value (V)](image)
Another tri-stimulus colour model is the CIE L*a*b model (Figure 3.5). The International Society of Illumination (Commission Internationale de l'Eclairage) has developed several similar colour models over the years, based on human perception, and the L*a*b* model is the one that seems to best represent how we think about hair colour. In this model, colour is measured on three axes that correspond to the trichromatic model of human perception [58].
Figure 3.5 An illustration of the CIE L*a*b colour space showing the three axes: L* (Black = 0 to White = 100), a* (-100 = green to +100 = red) and b* (-100 = blue to +100 = yellow).[57]

The lightness, or intensity, of a colour is measured on the ‘L*’ axis on a scale from 0 (being black) to 100 (being white). The colour is also measured on the ‘a*’ axis that gives a value from -100 (being green) to +100 (being red). The ‘b*’ axis measures colour from -100 (being blue) and +100 (being yellow). This model has the same benefit of the HSV model in that it corresponds to human perception of colour instead of a device-based model and has the additional benefit of giving a grid point for each specific colour [57] and this grid point allows for the mathematical comparison of colour. In addition, this model has been used for several other studies of human pigmentation [16, 17, 34, 59]. It should be noted that, theoretically, the a and b axes have no maximum or minimum values but this research has used the cut off points of +/- 100 because these represent the practical limit of the instrument and software used for colour measurement [60].

The comparison of colours in the CIE L*a*b* system often uses the delta (Δ) symbol for the ‘change’ or ‘difference’ in colour. ΔL* would be the difference between two L* values while Delta E* (ΔE*) is an Euclidian measure of the total distance between two CIE L*a*b* colour values and is calculated according to the following formula [55]:

$$\Delta E^* = [(\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2]^{1/2}$$

Figure 3.6 illustrates a difference of 5 units along the L*, a* and b* axes and one can see that the differences are discernable, but not obvious. It should be noted that the
colours in Figure 3.6 are approximate and only for illustration purposes, due to the differences in the way various computer screens and printers reproduce colour.

![Colours in Figure 3.6](image)

**Figure 3.6** An example of a 5 unit difference along the L*, a* and b* axis.

Now that a system of measurement has been defined, a method must now be determined. There are many ways to determine the colour of an object. Reflective spectrophotometry (RS) measures the wavelengths and amount of light reflected from an object and provides its own light source. This gives an accurate and consistent colour reading. Applications of RS include the measurement of pigments for paint and materials manufacturing and for determining food quality [61]. Colour may also be judged by comparing it to a colour chart (e.g. Munsell Book of Color) [55] which, while simple, involves a certain amount of subjectivity on the part of the observer.

Reflective spectrophotometry has been used to objectively measure skin pigmentation [8, 59] and the technique has also been used on hair in different ways and for various purposes including cosmetic science [16, 17], for the determination of hair composition and structure [19], for comparison of measurement methods [59] and in recent genetic studies [41, 52]. It has been found to be an objective and effective way to measure hair colour.
In 2000, Shriver and Parra used the CIE L*a*b* system with reflective spectrophotometry to measure the hair colour of 41 European-American individuals and 18 individuals of non-European ancestry, comparing the L* colour component to the Melanin Index [59]. They found a significant correlation between these systems of measurement and reported low variability in the hair colour of individuals of non-European ancestry. Naysmith et al. in 2004 used the CIE L*a*b* system to measure the hair colour of 50 individuals for genetic and chemical studies of red hair [52]. They found relationships between variations in the gene MC1R and measured hair colour (most strongly with b*), as well as with chemical studies. It should be noted that people with red hair were deliberately over-represented in this sample and volunteers ranged in age from 6 to 72 years old (median of 35). Both studies emphasised the necessity and demonstrated the feasibility of measuring pigmentation objectively and accurately.

3.1.3 Aims

In every investigation, ‘hair colour’ must first be defined by the researchers. What one investigation calls ‘blonde’ however, might be what another calls ‘light brown’ and this inhibits the reproducibility of experiments and the direct comparison of results.

The aim of the research presented here was to objectively measure human hair colour and examine the variation found in a population with European ancestry, over all three colour axes and to analyse the variation of the measured hair colours to determine if and how hair colours form describable groups and how these match with our intuitive descriptions of hair colour.
3.2 Materials and Methods

3.2.1 Subjects

Subject recruitment and sampling procedures were conducted with the approval of the Victoria University Human Research Ethics Committee (approval number HRETH06/156). In total, 202 subjects were included in this analysis. All volunteers recruited for the study had their natural hair colour at the time of sampling. Most volunteers were between the ages of 18 and 35 and were of European ancestry (194 subjects). The exceptions to these criteria include six individuals of non-European ancestry (one African and five South Asians) and two older European individuals with white hair. Subjects fitting these criteria were chosen to meet the study aims of examining natural variation in adult European hair colour, with non-European and mature-white haired individuals included for comparison purposes.

3.2.2 Sampling

Subjects were first given all project information and the following procedures carried out following their written consent. Subjects filled out a questionnaire that, in addition to confirming their age, ancestry and natural hair colour status, asked them to report how they saw their own hair and eye colour (eye colour data was not analysed for this project). Hair colour was reported to be one of; Black, Dark Brown, Light Brown, Blonde, Red or White. Colour assessment by an observer (the researcher taking the samples) was also recorded. The project information sheet, consent forms and questionnaire can be found in Appendix 3.

Macroscopic hair colour was measured by reflective spectrophotometry (RS) using a Minolta CR-300 Chroma Meter (Konica Minolta, North Ryde NSW, Australia). The instrument was calibrated using a white tile (supplied by the manufacturer) and a light source input setting of ‘D65’ which represents daylight without spectral highlights. The instrument was set to measure in the CIE L*a*b* format. Hair colour was measured on the left, right and back of the head and was measured five times at each spot (See Figure 3.7), the mean of the 15 measurements being the focus of subsequent statistical considerations.
Figure 3.7 The locations on the head where reflective spectrophotometry measurements were taken; 5 measurements were taken at each site.
3.2.3 Statistics

SPSS 15.0 for Windows (© SPSS Inc. 2006, Chicago IL) was used to analyze the colour data. Two separate methods were used for grouping the population; a cluster analysis and discriminant analysis. Measured hair colour (by RS) was broken into several groups using a two-step cluster analysis; performed using the Euclidian distance criterion or the log-likelihood probability approach and either not specifying the number of clusters to be formed or asking for a specific number of clusters, ranging from two to seven. Cluster analysis seeks to identify natural subgroups within a population by minimising within-group variation while maximising between-group variation. To characterise and evaluate the clusters a separate, discriminant analysis was then performed. Discriminant analysis commences from the perspective of there being a known number of population sub-groups. From the population, there are a number of individuals whose sub-group classification is known. The data from these individuals is analysed in an attempt to build a profile of sub-group membership and subsequently use this profile to classify new individuals (whose sub-group membership is otherwise unknown).

After the analysis, SPSS determines which variables are most important in discriminating the groups, and how good this model will be at predicting group membership of future individuals. It determines the predictive value of the model by removing each individual separately, reanalysing the data and then predicting the membership of that case. The percent correctly classified is reported.
3.3 Results

3.3.1 Reported Colours

Self-reported hair colours (SRC) were compared to the colours determined by the observer (observer-reported colour or ORC) (Table 3.1). 85.2% of individuals had the same hair colour reported by themselves and the observer. Where there was a difference, discrepancies were lighter or darker by one shade. The observer was more likely (53.6% of disagreeing observations) to see a darker shade. In one case, a self-reported Red colour was reported as Blonde by the observer.

Table 3.1 The self-reported hair colours (SRC) compared to the observer-reported hair colours (ORC) of individuals with European ancestry.

<table>
<thead>
<tr>
<th>Colour</th>
<th>Self Reported</th>
<th>Observer Reported</th>
</tr>
</thead>
<tbody>
<tr>
<td>Black</td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td>Dark Brown</td>
<td>85</td>
<td>71</td>
</tr>
<tr>
<td>Light Brown</td>
<td>50</td>
<td>62</td>
</tr>
<tr>
<td>Blonde</td>
<td>41</td>
<td>38</td>
</tr>
<tr>
<td>Red</td>
<td>14</td>
<td>13</td>
</tr>
<tr>
<td>White</td>
<td>2</td>
<td>2</td>
</tr>
</tbody>
</table>

3.3.2 Population Variation

To examine the hair colour variation amongst the European population, the L*, a* and b* values of the 196 European individuals, as measured by reflective spectrophotometry, were recorded and the data subjected to statistical analysis. The variability observed in the entire sample is illustrated in Figure 3.8. This distinctiveness of the non-European individuals (which are included and labelled as NE in Figure 3.8) and the individuals with white hair (W in Figure 3.8) can be seen in the highlighted areas. Individuals not of European ancestry have hair that is much darker with a* and b* values very close to zero.
3.3.3 Grouping the Population

The aim of the statistical analysis was to investigate the major components of the colour variation and to see whether hair colour falls into groups that would be meaningful for further research. What qualifies as a meaningful group is subjective and may depend on the intended use of the data; however in this case, meaningful groups would be those that separate colours into several distinct groups. For practical purposes, it would also be useful to have groupings that correspond well to colours people already use to report hair colour (e.g. Blonde).

The European sample was broken into groups using the Two-Step cluster analysis. The analysis was first conducted without specifying the number of clusters, using either the Euclidian distance approach (E) or the log-likelihood probability approach (LL). Next, the numbers of clusters were specified, from two to six for both approaches (LL2 to LL6 and E2 to E6) and a discriminant analysis was then performed. Using the clusters determined as a definitive population sub-group, the percentage of cases correctly classified was used to evaluate the clusters. A maximum of six clusters were analysed since there were six observer-reported categories and since discriminant scores decreased continuously with more clusters (data not shown).
The percentage of cases correctly classified in the discriminant analysis for the various methods of cluster determination is shown in Figure 3.9. When the observer-reported colour (ORC) groupings are used, only 70.4% of cases are correctly classified. Almost all other clusters have greater than 95% of cases correctly classified. Only E4 had less than 95.0% but was very close at 94.9%.

![Figure 3.9](image_url) Discriminant scores for each number of clusters and the method by which they were determined.

Using the Euclidian distance measure to determine clusters did not result in as useful groupings. In all cases, between 67% (for six clusters) and 93.9% (for two clusters) of the sample was in one cluster (data not shown). This is a problem because it results in one large cluster with only extreme individuals in several very small clusters.

The log-likelihood probability method of clustering produced several promising groupings (Table 3.2 to Table 3.4). The best result for the purposes of this study was obtained with five clusters (LL5). This has the highest discriminant score (99.5%) and the most appealing cluster characteristics (Table 3.2) which will be discussed later in more detail. Other numbers of clusters determined by log likelihood may be useful to different types of studies and have good discriminant values. For example, six clusters (LL6) would be useful when comparing to the six observer-reported groups discussed in Section 3.3.4 and in Figure 3.11, while two clusters (LL2) would be useful for dividing
a population into light and dark hair colours. Both of these analyses have good
discriminant scores (97.4% and 96.9%, respectively) and good group composition.

Table 3.2 Composition and cluster characteristics of five clusters determined by the log-
likelihood probability method (LL5)

<table>
<thead>
<tr>
<th>Cluster</th>
<th>N</th>
<th>% of Total</th>
<th>L* Mean</th>
<th>L* SD</th>
<th>a* Mean</th>
<th>a* SD</th>
<th>b* Mean</th>
<th>b* SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>8</td>
<td>4.1%</td>
<td>39.58</td>
<td>2.74</td>
<td>8.29</td>
<td>1.39</td>
<td>12.79</td>
<td>1.95</td>
</tr>
<tr>
<td>2</td>
<td>32</td>
<td>16.3%</td>
<td>46.80</td>
<td>5.38</td>
<td>4.42</td>
<td>0.65</td>
<td>11.19</td>
<td>2.01</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>1.0%</td>
<td>74.62</td>
<td>6.27</td>
<td>0.26</td>
<td>1.24</td>
<td>7.57</td>
<td>1.34</td>
</tr>
<tr>
<td>4</td>
<td>68</td>
<td>34.7%</td>
<td>35.46</td>
<td>2.44</td>
<td>3.22</td>
<td>0.62</td>
<td>4.82</td>
<td>2.26</td>
</tr>
<tr>
<td>5</td>
<td>86</td>
<td>43.9%</td>
<td>30.41</td>
<td>1.63</td>
<td>1.70</td>
<td>0.63</td>
<td>1.80</td>
<td>1.19</td>
</tr>
<tr>
<td>Total</td>
<td>196</td>
<td>100%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3.3 Composition and cluster characteristics of two clusters determined by the log-
likelihood probability method (LL2)

<table>
<thead>
<tr>
<th>Cluster</th>
<th>N</th>
<th>% of Total</th>
<th>L* Mean</th>
<th>L* SD</th>
<th>a* Mean</th>
<th>a* SD</th>
<th>b* Mean</th>
<th>b* SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>47</td>
<td>24.0%</td>
<td>45.88</td>
<td>8.40</td>
<td>4.87</td>
<td>1.96</td>
<td>10.99</td>
<td>2.33</td>
</tr>
<tr>
<td>2</td>
<td>149</td>
<td>76.0%</td>
<td>32.44</td>
<td>3.08</td>
<td>2.31</td>
<td>0.93</td>
<td>2.96</td>
<td>2.13</td>
</tr>
<tr>
<td>Total</td>
<td>196</td>
<td>100%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3.4 Composition and cluster characteristics of six clusters determined by the log-
likelihood probability method (LL6)

<table>
<thead>
<tr>
<th>Cluster</th>
<th>N</th>
<th>% of Total</th>
<th>L* Mean</th>
<th>L* SD</th>
<th>a* Mean</th>
<th>a* SD</th>
<th>b* Mean</th>
<th>b* SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>8</td>
<td>4.1%</td>
<td>39.58</td>
<td>2.74</td>
<td>8.29</td>
<td>1.39</td>
<td>12.79</td>
<td>1.95</td>
</tr>
<tr>
<td>2</td>
<td>27</td>
<td>13.8%</td>
<td>47.79</td>
<td>5.19</td>
<td>4.41</td>
<td>0.65</td>
<td>11.65</td>
<td>1.66</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>1.0%</td>
<td>74.62</td>
<td>6.27</td>
<td>0.26</td>
<td>1.24</td>
<td>7.57</td>
<td>1.34</td>
</tr>
<tr>
<td>4</td>
<td>27</td>
<td>13.8%</td>
<td>38.48</td>
<td>2.24</td>
<td>3.79</td>
<td>0.72</td>
<td>7.51</td>
<td>1.30</td>
</tr>
<tr>
<td>5</td>
<td>75</td>
<td>38.3%</td>
<td>30.14</td>
<td>1.54</td>
<td>1.58</td>
<td>0.55</td>
<td>1.73</td>
<td>1.12</td>
</tr>
<tr>
<td>6</td>
<td>57</td>
<td>29.1%</td>
<td>33.92</td>
<td>1.98</td>
<td>2.92</td>
<td>0.52</td>
<td>3.39</td>
<td>1.78</td>
</tr>
<tr>
<td>Total</td>
<td>196</td>
<td>100%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
During analysis, canonical discriminant functions are determined. These unstandardised functions are used to make classifications in the discriminant analysis and the standardised function coefficients are used to compare the relative importance of each function and each variable in making classifications. For LL5, three discriminant functions were determined; F1 = -7.359 + 0.089L* + 0.953a* + 0.518b* which contributes 66.6% of the information for discriminating clusters, F2 = -9.759 + 0.373L*- 0.822a* – 0.233b* with 32.3% of the information and F3 = -4.802 + 0.140L* + 0.826a* – 0.534b* with 1.1% of the information. In Figure 3.10, the results of the first two functions are used to illustrate the clusters.

Figure 3.10 Five clusters determined by the log-likelihood probability method (LL5) separated according to the results of the first two significant discriminant functions. F1 = -7.359 + 0.089L* + 0.953a* + 0.518b* and F2 = -9.759 + 0.373L*- 0.822a* – 0.233b*
3.3.4 Comparing Clusters to the Observer-Reported Colours

It appears that the clusters determined by various methods show similarity to hair colours as perceived by an observer. Looking at the LL5 clusters in Figure 3.10 again, the clusters could be referred to as Red, Fair, White, Medium and Dark (for clusters 1 to 5, respectively) by both their L*a*b* values (Table 3.2) and the observer reported colours they contain (Table 3.5). The names are arbitrary; however they correspond to how people tend to refer to hair colour.

Table 3.5 Observer reported colours found in the LL5 clusters and descriptions of individuals compared to other ORC group members.

<table>
<thead>
<tr>
<th>LL5 Cluster</th>
<th>Possible Cluster Name</th>
<th>Observer Reported Colours Present</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>Red</td>
<td>8 Red (highest a* and b* values)</td>
</tr>
<tr>
<td>C2</td>
<td>Fair</td>
<td>29 Blonde</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3 Red (less a* and b* than C1 )</td>
</tr>
<tr>
<td>C3</td>
<td>White</td>
<td>2 White</td>
</tr>
<tr>
<td>C4</td>
<td>Medium</td>
<td>10 Blonde (darker and less a* and b* than C2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>41 Light Brown</td>
</tr>
<tr>
<td></td>
<td></td>
<td>15 Dark Brown (lighter and higher a* and b* values than C5)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2 Red (less a* and b* than C1, darker than C2)</td>
</tr>
<tr>
<td>C5</td>
<td>Dark</td>
<td>16 Light Brown (darker and less a* and b* than C4)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>62 Dark Brown</td>
</tr>
<tr>
<td></td>
<td></td>
<td>8 Black</td>
</tr>
</tbody>
</table>

While the five clusters determined in LL5 are the best for the purposes of this project, it is worthwhile illustrating a direct comparison between the six observer-reported categories and six clusters produced by SPSS. Figure 3.11A shows the six clusters as determined by their reflective spectrophotometric measurements (LL6), separated according to the discriminant functions while Figure 3.11B shows the same individuals grouped into their observer-reported categories. The lack of separation using observer-reported categories is obvious and reflects the low mathematical discrimination ability of these categories. It should be noted that clusters are numbered independently in each analysis; Cluster 1 in LL5 does not necessarily contain any or all of the same individuals as Cluster 1 in LL6.
Figure 3.11  The two most important functions used to discriminate the six clusters as determined by log-likelihood (LL6) were applied to the L*a*b* data and grouped by Cluster (A) and by observer-reported colour (B). \( F1 = -8.558 + 0.101L^* + 0.972a^* + 0.435b^* \) and \( F2 = -9.671 + 0.376L^* - 0.908a^* - 0.222b^* \)
3.4 Discussion

The presented data show that when given a limited choice of colours, self-reported hair colours and those reported by an observer are fairly consistent; however, these categories are not necessarily the best way to categorise hair colour for quantitative research. While hair colour can be perceived as a continuum (as seen in Figure 3.8), it can be divided into categories or clusters based on spectrophotometric measurements in the CIE L*a*b* colour space and these clusters can be well discriminated from each other. The discrepancies between self-reported hair colour and observer-reported hair colour and the poor separation of observer-reported colour groups compared to those defined by clustering analysis emphasise the importance for objective measurement of individuals to be included in research studies. For repeatability and validity of studies, phenotypes must be defined as well as possible and the objective measurement methods and analysis strategies presented here may be of assistance in this regard. The reduction of three colour components to one or two discriminant functions makes the examination and classification of individuals easier to represent.

It is also interesting to note that, while not exact, clustered groupings follow the same pattern that self-and observer reported colours do. How someone decides to report a hair colour may be an artefact of the arbitrary naming of colours or of personal bias or it may also be due to human perception of colour, which corresponds to the CIE L*a*b* system of measurement or it may be all of the above. The human eye is a complicated structure with retinal receptors (rods and cones) that are sensitive to changes in lightness and colour [55] as discussed in Section 3.1.1. This system of human trichromatic vision may be why similar groups are defined by both the human reporting of colour and by the spectrophotometry and clustering method described here. The lack of sensitivity in biological perception may contribute to the variability found. As mentioned previously, one unit on the L*a*b* scale is defined as the limit of human discrimination between colours [57]. The difference between members of different clusters can be less than this in one or more of the three colour axes and may contribute to the inconsistency between reported colour groups and clusters determined by spectrophotometric data and clustering algorithms.

The particular clustering algorithms and program options used will depend on the intended use of the data. For example, a medical genetic association study may require
a Fair/Dark division, where the prediction of hair colour in a forensic study may require more specific groupings. This study has illustrated the possibilities by using a two-step cluster analysis in SPSS; however there are many other programs and clustering algorithms available. Having objectively measured colours allows researchers to group individuals in clearly defined ways and to change these groups as the study demands without subjectivity or collecting additional data.

It is acknowledged that the exact functions and cluster details may change as larger samples are analysed. In this study, the composition of clusters changed when the sample size was increased from 134 to 196 (See Vaughn et al. 2008 in Appendix 4). LL2 and LL6 retained their high discriminant scores and good group composition; however E4 did not retain its high discriminant score and was replaced by LL5. It is expected that a sample size would eventually become sufficiently large to resist major changes. Using smaller sample sizes with colour measurement data, the researcher has the options of changing clusters for each sample-set or defining groups based on an initial sample-set and assigning new individuals according to the associated discriminant functions.

As mentioned, different clustering results may be appropriate for different types of studies. For future considerations in this research, the LL5 clusters are the most appealing. This grouping conveniently divides the non-red (blonde-black) spectrum into three large groups and recognises those with Red and White hair as separate. This corresponds nicely with the study aims of studying non-red colours.

Reflective spectrophotometry, while accurate, may be inconvenient for measurements on very large numbers of people or for investigators. Chapter 4 will investigate the use of digital images for measuring and classifying colour.
Chapter 4  A comparison of Hair Colour Measurement by Digital Image Analysis with Reflective Spectrophotometry

4.1 Introduction

4.1.1 Digital Photography and Digital Colour Measurement

Traditional photography captures images by exposing light sensitive silver compounds while digital photography captures images by a charge coupled device (CCD) array which transforms the absorbed light energy into a digital image [62] that can be viewed on a computer screen. Figure 4.1 illustrates the two processes of image production.

![Figure 4.1 Image formation by silver halide and CCD sensors [62]](image)

The reproduction of colour in photography is a complex issue and while no reproduction of colour is identical with the object being photographed, modern imaging

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3 An earlier version of this experiment was published in Forensic Science International. Vaughn et al. 2009:183 Pages 97-101. Please see Appendix 4
systems are able to reproduce colour very well [62]. Digital images are usually displayed in the RGB colour space (Discussed in Section 3.1.2) on a computer monitor or similar device. The RGB colour system is composed of three values (Red, Green and Blue) that are measured from 0 to 255. Therefore a colour 0,0,0 is pure black while 255, 255, 255 is pure white and various colours or hues are made of combinations of red, green and blue [57]. Figure 4.2 illustrates the colours and related values of the RGB model.

![Figure 4.2 Colours and values within the RGB model](image)

Colours can be measured by various software applications by reporting the intensities of the red, green and blue in the image. Conversion to other colour spaces is then a matter of mathematical algorithms [63]. Different colour spaces are not equivalent to one another but images can be measured in various spaces or the values converted. For this experiment, we are concerned with converting RGB to the CIEL*a*b* space discussed in 3.1.2. The formulas for converting RGB to L*a*b* from Herbin et al. [63] are:
\[ L^* = 25 \times (100 \times \frac{Y}{Y_0})^{1/3} - 16 \]
\[ a^* = 500 \times [(\frac{X}{X_0})^{1/3} - (\frac{Y}{Y_0})^{1/3}] \]
\[ b^* = 200 \times [(\frac{Y}{Y_0})^{1/3} - (\frac{Z}{Z_0})^{1/3}] \]

where XYZ is the CIE system defined by:

\[ X = 0.618 \times R + 0.177 \times G + 0.205 \times B \]
\[ Y = 0.299 \times R + 0.587 \times G + 0.114 \times B \]
\[ Z = 0.000 \times R + 0.056 \times G + 0.944 \times B \]

and \(X_0, Y_0, Z_0\), are the values of a specific white reference.

While reflective spectrophotometry is an established method for measuring hair colour at the macroscopic level, it can be cumbersome to use on a large number of individuals. This study investigates the use of digital photographs to measure hair colour and compares its use to the reflective spectrophotometry standard. An understanding of the accuracy of the colour determination by these methods is of relevance when undertaking specific investigations, such as those on the genetics of hair colour. Measurement of hair colour by digital image analysis may also be of assistance in cases where a photograph is the only evidence of hair colour available (e.g. surveillance).

Digital image analysis has previously been used to measure hair colour in forensic science; however it has generally been used to measure the colour during the microscopic examination of single hairs and not the macroscopic colour of hair on the head [24]. This has been done to assist the traditional forensic hair examination, with the purpose of attributing a hair to a specific donor. Digital images have been used extensively to measure the colour of many other objects, such as vegetables, soil and chocolate [64-66], and have been used to measure skin colour [63, 67, 68]. Herbin et al [63] found that the colour of skin lesions measured from 12 digitised images of colour slides was highly correlated with values obtained by reflective spectrophotometry, with correlation coefficients of 0.97, 0.99 and 0.98 for \(L^*\), \(a^*\) and \(b^*\) of the CIE \(L^*a^*b^*\) model (see Figure 3.5) respectively. Takiwaki et al. [67] achieved similarly good results with eight videomicroscopic images of skin.

When measuring colour from digital images, it is important to note that digital images are most often displayed and measured in the Red-Green-Blue (RGB) colour space of the computer monitor. There are various programs that may be used to measure this
colour and then to convert it into the colour space required (CIE L*a*b* in this case) using standard mathematical algorithms [63]. This study considers two such programs.

4.1.2 **Aims**

The aim of this study is to compare the results obtained from measuring the colour from digital images (DI) of hair colour (at the macroscopic level) to the measurements of the same individuals using reflective spectrophotometry (RS) as a standard, to see how the measurements correspond. This will help determine to what extent digital image analysis can be used instead of reflective spectrophotometry and, if so, in what situations.
4.2 Materials and Methods

4.2.1 Subjects

Described in 3.2.1

4.2.2 Sampling

In addition to the reported colours and reflective spectrophotometry measurements described in Section 3.2.2, digital photographs were taken of the scalp hair. Digital images were taken of the left, right and back of the head with a CASIO 4.0 mega pixel camera (model QV-R40) with an image size of 640x480 pixels and with picture quality set to ‘fine’. On ‘collection’ mode (for taking close-up photographs), the flash was turned off, the white balance tool was set to ‘sunlight’ and all other settings were left at the default. Each time a set of images was to be taken under a different light source (on a different day, different time of day, different room, etc.) a piece of new white paper was photographed in the same position as the head of the subject (three sides) and measured using the Minolta CR-300. This ‘white image’ was later used to correct colour for light source. All images were taken at a distance of 15cm in the same locations as the reflective spectrophotometry readings (See Figure 4.3 and compare to Figure 3.7).

![Sample photographs](image-url)

**Figure 4.3** Sample photographs before cropping and analysis
4.2.3 Digital Image Preparation and Analysis

Software packages used included:

- JASC Paint Shop Photo Album 4.0 by Jasc Inc.
- IrfanView 3.92 © 1996-2004 Irfan Skijan; Vienna University of Technology
- MATLAB 4.0 © 1994-2006 MathWorks with the Image Processing Toolbox and modules written to convert RGB colour measurements to L*a*b measurements [64]
- V++ Version 4.0 ©1990-2002 Digital Optics Ltd. New Zealand, with modules written to convert RGB colour measurements to L*a*b measurements [64, 69]
- ImageJ ©1997-2006 by W.S. Rasband at the US National Institutes of Health, with the RGB-to-Lab conversion plugin installed [70].

Each digital image of the hair was cropped to 100x100 pixels in the area closest to the centre of the image (where the RS measurements were taken) with JASC Photo Album, Adobe Photoshop or IrfanView as available. Care was taken to avoid obvious shadows or spectral highlights that result from the curvature of the skull.

Initially, images were analysed using MATLAB to convert the RGB colour measurements to L*a*b* and to determine the mean colour values for each image; however, this program is rather difficult and tedious for a new user. The RGB to L*a*b* conversion modules written for MATLAB were instead used within V++ [69], a more user-friendly program, which results in identical values as those determined by MATLAB. Therefore, V++ was used to analyse all subsequent images. The computer code for the RGB2LAB conversion module can be found in Appendix 2.

For each image, the L*, a* and b* values were calculated. While a reflective spectrophotometer has its own, consistent light source and can be calibrated to a white tile (as described in Section 3.2.2), a digital camera cannot be calibrated before the photograph is taken and since hair is shiny and reflects a flash, no consistent light
source was available. Images of the same colour taken under different light sources will appear different [63], so an effort was made to correct this colour in a manner similar to a previous study using the RGB colour space [63], with the understanding that perfect correction is unlikely and that the greatest difficulty in correcting the colours (a* and b* axes) will be found when the lightness of the image differs. The new white paper, mentioned above, when measured with the Minolta Chroma Meter CR-300 was found to have L*, a* and b* values of 93.4, 1.16 and -2.06 respectively. Using this as a reference point, L*, a* and b* colour points were added or subtracted depending on how much the white reference paper deviated from its Minolta CR-300 determined values under the various sampling situations. For example, if \( L_i \) is the L value of the colour image, \( L_w \) is the L value of the white image and \( L_c \) is the L value of the corrected image, then: \( L_c = (93.4 - L_w) + L_i \).

### 4.2.4 Correction

To examine the accuracy of the correction formula, images were taken of white, pink and blue paper under four different light sources i) fluorescent light ii) fluorescent light and natural sunlight iii) incandescent light and natural sunlight and iv) bright natural sunlight. The images of the paper were cropped to 100x100 pixels and analysed for colour with V++. The pink and blue colours were corrected using the white values obtained from V++, using the above formula and compared to the RS measurements (true colour) of the paper.

### 4.2.5 Program Comparison

To examine whether another program is better able to match the RS values through its algorithm or data handling, the MATLAB/V++ module for RGB to L*a*b* colour conversion was compared to that of the ImageJ program. Values were calculated for the pink, blue and white paper images and corrected using the white paper values calculated from Image J and with the above formula. The corrected ImageJ values were compared to the RS measurements and to the V++ values.
4.2.6 **Statistics**

SPSS 15.0 for Windows (© SPSS Inc. 2006, Chicago IL) was used to analyze the colour data. Basic statistical tests were performed, including a One-Sample T-test (two tailed) for a difference in mean, a Paired T-test (two tailed) and an F-test for a difference in variance [71]. For details on the cluster analysis using reflective spectrophotometric data, please see Section 3.2.3 [72].

Delta E* ($\Delta E^*$) is an Euclidian measure of the total distance between two CIE L*a*b* colour values and is calculated according to the following formula [55]:

$$\Delta E^* = [(\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2]^{\frac{1}{2}}$$
4.3 Results

4.3.1 Colour Correction

When comparing the colour mean values of pink paper measured by digital image analysis to the reflective spectrophotometric value, it was found that the DI $a^*$ component values have means that are significantly lower than the RS measurements, whether the correction formula is used or not. Using a One sample T-test, there was a 95% confidence interval of -1.51 to -5.96 for the difference between the uncorrected DI measurement for $a^*$ and the RS measurement and from -0.89 to -8.27 for the difference between the corrected DI measurement for $a^*$ and the RS measurement, thus neither of these CI ranges included zero, indicating a significant difference. There was no significant difference between the RS and DI means for the $L^*$ or $b^*$ values. For the blue paper, only the corrected $L^*$ value was significantly different from the RS measurements with a 95% confidence interval (One sample T-test) of 3.78 to 9.90.

Although not all of the means were significantly different, it was noted that for the $L^*$ and $b^*$ values of Pink and Blue paper (four out of six measurements) the confidence intervals for corrected digital measurements were narrower than those for the uncorrected formulas, and so an F-test for the equality of variance was conducted (between the uncorrected and corrected values for pink and blue paper). The results indicate that the variance for $L^*$ of pink paper and the $L^*$ and $b^*$ components of blue paper are significantly smaller using the correction formulas (F-test for variance, p-value <0.05). It was decided to use the correction formula for further experiments due to the closer means (though not significant) and the significantly smaller variances.

4.3.2 Program Comparison

To test whether the colours obtained by the programs V++ and Image J are any different, a Paired T-test of the means was used for the corrected $L^*$, $a^*$ and $b^*$ values from each program. The only significant difference (p value = 0.002) was that the $a^*$ value for the blue paper was slightly higher in ImageJ (-5.19) than in V++ (-5.99).

The variance of the measurements as determined by the two programs was also compared. There was no significant difference (F-test for variance; p-values > 0.1) between the variance of colour measurements as determined by V++ and ImageJ. It can
be concluded that the L*a*b* colour measurement functions of ImageJ and V++ can be used interchangeably.

4.3.3 Intra-individual Comparison

To test the consistency of the digital image colour measurements, 10 sets of digital images were taken of one individual (three photos for each set taken of the right, left and back of the head as seen in Figure 3.7). The mean L*, a* and b* of each set (of three images) was calculated and a one-way ANOVA was used to determine that these means were not significantly different over the 10 sets. This indicates the low amount of variation found across multiple measurements taken of one individual under the same conditions.

4.3.4 Digital Image Comparison to Reflective Spectrophotometry

After determining that the correction formulas produce more consistent results and that two programs selected to analyse the digital images are almost the same, the entire European sample (196 individuals) was analysed to determine how digital image (DI) colour analysis compares to the colours as determined by reflective spectrophotometry (RS). The mean ΔE* value for the difference between RS and DI colour was 34.16 units with a standard deviation of 9.10 units. However; this measurement does not give information about which components are different and therefore, each colour axis was examined separately.

Initially, the digital image colour measurements were compared directly to the RS measurements for the L*, a* and b* axes. A correlation was found along all three axes, with Pearson Correlation Coefficients of 0.675, 0.679 and 0.537 for L*, a* and b* respectively (p-values = 0.000). This is a significant correlation, but of only moderate strength. The coefficient of determination (R²) indicates that only 45.6%, 46.1% and 28.8% of the variation (for L*, a* and b*, respectively) can be explained by the relationship between the RS and DI measurement. Figure 4.4 illustrates this relationship.
i)

\[ R^2 = 0.4555 \]

ii)

\[ R^2 = 0.4607 \]
iii)

**Figure 4.4** The relationship between RS and DI measurements for i) L* ii) a* and iii) b*

To see if there is a pattern in the differences between DI and RS measurements, a Paired T-test for the difference between the means obtained by DI and RS was performed. Means are significantly overestimated by digital image analysis for all three colour components. The L* value is significantly higher by an average of 31.28 units, the a* value higher by an average of 3.32 units and the b* value higher by an average of 5.68 units (p-values = 0.000).

To investigate whether the overestimation error was uniform across the sample, or if light or dark hair is more prone to overestimation, the difference between the DI and RS measurements was compared to the original colour (RS measurement). The difference in L* showed no significant correlation, indicating that the amount of overestimation is not related to the original hair colour for this component; however, there was a low amount of correlation (R = 0.341) between the overestimation of the DI a* and the RS a* values (p-value = 0.000) and between the overestimation of the DI b* and the RS b* values (R = 0.261, p-value = 0.000) indicating that there is a slight tendency for the
difference in a* values to become greater as hair becomes more red and for the difference in b* values to become greater as hair becomes more yellow.

As the separate colour components (L*, a* and b*) measured by RS and DI do not correspond well with each other, it was investigated whether, when combined, they would place individuals into the same colour categories as those determined by RS measurements.

Chapter 3 showed that the RS measurements can be used to break this same sample set into defined groups using a Two-Step Cluster analysis. These groups were then evaluated using a discriminant analysis to classify cases into the correct groups. In Chapter 3, the best results were obtained for an analysis resulting in five clusters using a log-likelihood probability method (LL5) with a discriminant score of 99.5%. Other groupings may have some use, such as LL2 or LL6 and each of these obtained classification scores above 95%. This was better than the 70.4% discrimination obtained by using observer-reported clusters.

For the present study, each individual was assigned to a cluster based on their RS data (for each of the three analyses). These clusters were then re-evaluated, using their DI data and the percent of cases correctly classified is reported in Table 4.1. Table 4.1 also shows that the percent of cases correctly classified in the six observer-reported colour groups (52.0%) was lower than for any of the RS analyses.

<table>
<thead>
<tr>
<th>Analysis</th>
<th>Number of Clusters</th>
<th>Percent Correctly Classified by Digital Image Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>LL2</td>
<td>2</td>
<td>82.1%</td>
</tr>
<tr>
<td>LL5</td>
<td>5</td>
<td>63.8%</td>
</tr>
<tr>
<td>LL6</td>
<td>6</td>
<td>58.2%</td>
</tr>
<tr>
<td>ORC</td>
<td>6</td>
<td>52.0%</td>
</tr>
</tbody>
</table>

*Table 4.1* Percent of cases correctly classified by a discriminant analysis on Clusters determined by RS or the observer reported colour group (ORC), using DI data.
Figure 4.5 illustrates an example of the use of discriminant functions in classifying individuals (using the results from LL5). The discriminant functions determined by RS data were $F_1 = -7.359 + 0.089L^* + 0.953a^* + 0.518b^*$, $F_2 = -9.759 + 0.373L^* - 0.822a^* - 0.233b^*$ and $F_3 = -4.802 + 0.140L^* + 0.826a^* - 0.534b^*$. The first two functions were used to plot individuals onto a grid using first their RS measurements (Figure 4.4A) and then their DI measurements (Figure 4.4B). Figure 4.4A shows the good separation of the five clusters determined in LL5 while Figure 4.4B shows the groups merging into one another.
Figure 4.5 RS Clusters with RS data (A) compared to DI data (using RS discriminant functions) (B) for LL5. Clusters are numbered 1 to 5.
4.4 Discussion

The presented data have shown that although measuring colour from digital images is convenient, the colours measured and classified do not correspond very well to those colours measured by reflective spectrophotometry and would have limited use in studies requiring accurate measurement of hair colour, such as genetic association studies. They also do not provide an accurate description of hair colour for crime investigators to utilise as visual identifiers or in description announcements. Some use might be found in distinguishing between a small number of clusters with digital images, as in the two clusters of Analysis 1 (which showed 82.1% of cases correctly classified); however, as the number of clusters or groups increases, there is a lower correspondence. As shown in Chapter 3, observer-reported colours were not well discriminated by RS data (70.4% of cases correctly classified) and were more poorly discriminated by using digital image analysis (52.0% of cases correctly classified). These results support the assertion that objectively defined and measured colours, rather than observer or self-reported colours are necessary to classify individuals for further study and at this time, this is best achieved through reflective spectrophotometric measurement.

It should be noted that the images in this study were all taken with the same camera under the specific conditions of a well-lit room and from a close, specified and identical distance and all treated in the same way. Any additional variability in conditions would contribute to less similarity between the digital image analysis and a reflective spectrophotometry standard. If variability could be reduced, through improvement in photographic conditions or through more accurate correction for varying conditions, results may improve. The act of photographing an object introduces error in colour reproduction due to the nature of the camera and image production and is an approximation of the visual field (whether digital or with film)[62]. Different models of camera may have different abilities to reproduce colour accurately and may improve results. The process of measuring RGB values and converting RGB measurements to CIE L*a*b* colour space may also result in the loss of some colour information.

It may also be that there are components of hair colour that are omitted when measuring colour directly from a digital photograph. This may also part of reason for the higher correlations found by Herbin et al.[63] and Takiwaki et al.[67] in measuring skin colour by digital image analysis. For example, feature such as hair texture, thickness or lustre
may have a different effect on the analysis of colour from a digital image than they have on reflective spectrophotometry or on human perception. These features would have a lesser, or a different, effect than when measuring skin colour. With these things in mind, the improvement of digital cameras and further research into the digital photography of hair would assist in developing methods that provide a more accurate measurement and classification of hair colour that would be of use to researchers, investigators and forensic scientists.

Chapter 5 will introduce microscopic measurements of hair colour and morphology.
Chapter 5  Microscopic Examination of Hair Colour and Colour Morphology

5.1  Introduction

5.1.1  Microscopic Hair Structure

Hair is commonly encountered as physical evidence at crime scenes [21]. Traditional forensic hair examination involves examining the specimen under a microscope and observing characteristics such as colour, length and diameter of the hair, as well as the distribution of pigment granules and other morphological features of the hair that may assist in determining the source of the hair [21-24].

Hair is an appendage of the skin [21, 73] and consists of a hair root and hair shaft that grow from a hair follicle (Please see Figure 5.1). For a traditional forensic hair examination, as well as for the measurement of hair colour, the hair shaft is the most important part and is covered in more detail in Figure 5.2.

![Figure 5.1 A cross section of a hair follicle.](image)

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4 Parts of this chapter have been published in Microscopy and Microanalysis. Vaughn et al. 2009:15 Pages 189-193. Please see Appendix 4
Figure 5.2 Illustrates the microscopic features of the hair shaft. a) Cartoon of the hair shaft with cut-through to reveal inner aspects. Co, cortex; Cu, cuticle; Md, medulla. A macrofibril (300 nm) of single cortical cell is revealed with a constituent microfibril (MF; 7–10 nm). b) Whole mount of a scalp hair shaft with focal plane on cuticle (Cu, left) and medulla (Md; middle). c) Transverse section of a fine hair shaft revealing a multilayered cuticle (Cu) encompassing a tightly packed cortex (Co) containing many dark melanin granules. From Tobin et al. 2006 [74].

5.1.2 Cellular Biology of Pigmentation

Melanocytes are dendritic, neural-crest derived cells found in the epidermis, hair follicle and iris [4]. Melanocytes in the hair follicle, specifically melanocytes in the proximal hair bulb [6], produce melanin. Melanin is produced in cellular organelles called melanosomes; these pigment granules are then secreted and taken up by the keratinocytes that make up the growing, pigmented hair shaft [4, 6]. Melanosomes are cellular organelles (similar to lysosomes) that package the melanin and have a role in its distribution [4, 6].

Once melanin is formed in the hair bulb it is passed to surrounding keratinocytes that will be incorporated into the hair shaft [4, 73]. Unlike skin, which is continuously pigmented, pigmentation of hair only occurs during the hair’s active growth stage (called the anagen stage) [6]. The mechanism for this transfer is uncertain. Several hypotheses have been proposed [6, 75]; the most likely method is by exocytosis of the melanin granules and uptake by the keratinocytes, the other possible mechanisms include cytophagocytosis of the melanocytic dendrite, fusion of the plasma membrane...
between the melanocyte and keratinocyte or membrane vesicles that are released and are ingested by the keratinocytes.

Eumelanin (the black/brown pigment) is produced in ellipsoidal melanosomes with a fibrillar matrix, while pheomelanin (the red/yellow pigment) is produced in smaller, spherical melanosomes with a vesiculoglobular matrix and irregular melanin deposition [4, 6, 51]. The switch between eumelanin and pheomelanin production is controlled by the melanocortin-1 receptor (MC1R) and proopiomelanocortin (POMC) ligand pathway [4, 14].

The shape of the melanosomes and the packaging of melanin granules may also influence colour of skin, eyes and hair by scattering light in different ways [4, 73]. In addition, darker pigmentation of skin, and possibly of hair, is associated with higher numbers of melanosomes, although the number of melanocytes remains constant [4, 51]. Melanocytes in black hair follicles contain the greatest number of melanosomes (which are eumelanosomes), while the melanosomes in brown hair bulbs are smaller and those in blonde hair are very poorly melanised [6]. While melanin in the epidermal keratinocytes of skin is almost completely degraded over time, this does not occur in hair; melanin remains “stored” in the hair shaft [6]. This is thought to be part of the mechanism that allows for situations such as dark hair and fair skin.

5.1.3 Microscopic Hair Colour Measurement

In an attempt to assist with the classification of hair colours in a forensic examination, Bednarek [24] illustrated one way of measuring hair colour at the microscopic level; calculating the Red-Green-Blue (RGB) values of a digital image of the hair [24]. The average RGB values for ‘Blonde’ and ‘Brown’ hair were determined.

Recent research has also been conducted on the measurement of hair colour at the microscopic level for determining the contributor of a hair for identification purposes [27]. By using a discriminant analysis to assess colour measurements, Brooks et al. [27] were able to correctly assign a hair to a particular individual between 58% and 68% of the time (depending on what colour model was used), based on known microscopic colour measurements of the individuals’ hair [27].
5.1.4 **Aims**

The aim of this experiment was to determine the association between colours measured at the microscopic level with colour measured at the macroscopic level and to evaluate two morphological properties of hair – thickness and pigment pattern.

Since the macroscopic hair colour is made up of hundreds or thousands of strands of slightly different colours, it is important to know if the microscopic colour of a strand can indicate what the head of hair might look like and vice versa. This would be useful if a strand of hair is found as evidence or for the prediction of both macroscopic and microscopic colour from DNA. Examining morphological properties of the hair, such as thickness [53] and pigmentation patterns, and their associations with colour provides more information on the physical basis of hair for the description of phenotypes.
5.2 **Materials and Methods**

5.2.1 **Subjects and Sampling**

From the 202 subjects described in Section 3.2.1, 36 were also included in the microscopic analysis, a number restricted by available samples and time available to use the microscopic equipment. This included two individuals with white hair, three individuals of non-European ancestry and 31 individuals of European ancestry between the ages of 18-35, all with their natural hair colour. Photographs of these individuals’ hair can be seen in Appendix 6.

In addition to samples and measurements described in Sections 3.2.2 and 4.2.2, several full-length strands of hair were cut from the scalp of these volunteers.

5.2.2 **Preparation of Hairs for Microscopic Analysis**

Hairs were mounted on Livingstone Premium glass slides (76.2 x 25.4 mm pathology grade) with Entellan mounting medium (Merck) and covered with Marienfeld cover slips (22 x 60 mm No. 1). This medium has a refractive index of 1.5, which is required for proper viewing under the conditions described in Section 5.2.3. The area to be examined was marked on the slide. Three areas on each of three hairs were examined for each individual and the mean of these measurements was used for most analyses. A hair prepared for microscopic analysis can be seen in Figure 5.3.

![Hair prepared for microscopic analysis](image)

*Figure 5.3 Hair prepared for microscopic analysis.*
5.2.3 Generation and Collection of Hair Colour Data

Microscope and Image Collection software was generously made available for use at the Australian Federal Police Laboratories in Weston, ACT. Protocols for microscopy and montage were developed and optimised by Elizabeth Brooks for her MSc thesis [27]. Protocols for colour measurement were developed by Bruce Comber (AFP), Elizabeth Brooks (AFP) and myself with some source code for the conversion of RGB to L*a*b* provided by Alasdair McAndrew and Zhenliang Shen (Victoria University, School of Computer Science and Mathematics).

Equipment and Software

*Leica DMLB* Microscope with a 50x oil immersion lens (500x magnification; field of 276µm), using 10.5V illumination (daylight equivalent) and with DLF (daylight filter) on.

*Olympus DP70* 12.5 mega pixel digital camera (Olympus Optical Co. Ltd. 2003) and *Image Management System* which consists of the DP Controller (Version 2.1.1.183) and DP Manager (Version 2.1.1.163). An image size of 1360 x 1024 pixels was used with ISO 200 sensitivity, exposure time set to ‘automatic’, 1% (of the field) contrast correction and all images saved as TIF files. See Figure 5.4

*Automontage-Pro 5.02.0096* software (by Syncroscopy; a division of Synoptics Group, UK) was used to montage images of the middle third of the hair (the area containing the greatest amount of pigmentation information) [27]. Hair thickness was estimated by the depth of the optical field. See Figure 5.5

*V++ Version 4.0* (by Digital Optics Ltd. New Zealand, ©1990-2002) with modules written to convert RGB colour measurements to L*a*b* measurements was used to extract the correct areas of the microscopic images and to measure the colour of the extracted area [64, 69].

*Adobe Photoshop CS2* (by Adobe Systems Inc., San Jose, California, USA © 2005)

The procedure for the collection of the montage images is as follows. On the back of each slide, three points along the hair were marked (Figure 5.3). The hair was found with a 20x lens and the camera adjusted so the hair ran horizontally across the screen, then the magnification was increased using the 50x lens.

The 1% contrast box was dragged to a position so that half of the box was over the hair and half was over the background. The white balance tool was used by clicking on the background.

The depth of the hair was found by noting the number on the fine focus knob; first when the pigmentary features came into focus (or for very light hairs, when the cortical scales disappeared from focus) and next when the pigmentary features ended (or the cortical scales returned to focus). E.g. 40 – 55 = 15µm depth.

The depth of the field was divided by three. This gives the number of optical planes required for the middle third of the hair. From the previous example; 15 ÷ 3 = 5 optical planes that start at 45.

Images were captured for the middle third of the hair (e.g. at 45, 46, 47, 48 and 49) and saved to file. These are the ‘source images’. The source images were opened in Automontage-Pro (Figure 5.6) using the following settings: ‘align’ source images’, ‘scan montage’, and ‘save the montage to file’.

The montaged image was opened in V++ and was ‘extracted’ by highlighting the hair and removing the image background. Edge artefacts (resulting from light refraction) were removed from the image. If hairs contained a black medulla (an artefact of the slide mounting medium), this was also removed with Adobe Photoshop by cutting the top part of the image and moving it over the medulla (Figure 5.7). The new ‘extracted montage’ image was saved to a separate folder (Figure 5.8 and Figure 5.9)

In V++, the folder with all extracted montages was analysed using the rgb2lab function. Since all images were taken under identical lighting conditions, no corrections were made to the resulting L*a*b values. The results were copied into an Excel spreadsheet for analysis.
**Figure 5.4** Screenshot of the Olympus DP Manager used to take images

**Figure 5.5** Screenshot of the Automontage software
Figure 5.6 – Montaged image

Figure 5.7 Image of a black medulla that was later removed with Adobe Photoshop

Figure 5.8 – Extracted image of a ‘Light Brown’ hair

Figure 5.9 – Extracted image of a ‘Blonde’ hair
5.2.4 **Generation and Collection of Hair Thickness Data**

Hair thickness was estimated by the depth of the optical field, as described in Section 5.2.3. The thickness of the hair may have an effect on the colour by affecting how light travels through the hair and reflects from the pigment.

5.2.5 **Pigmentation Pattern**

The montage image produced in Section 5.2.3 was transformed into a monochrome image and the standard deviation of image intensity was measured. This measurement gives an estimate of how variable the density of the pigment granules is, which may also have an effect on the colour by affecting how light travels through the hair and reflects from the pigment.
5.3 Results

5.3.1 A Comparison of Microscopic Colour to Reflective Spectrophotometry

When comparing microscopic (Mc) colour L*, a* and b* measurements directly to those determined by RS, regression analysis found that a* and b* measurements fail to show any significant linear relationship between RS and Mc (p-values = 0.317 and 0.288, respectively). RS and Mc L* show a moderate, significant linear relationship with $R = 0.631$ ($R^2 = 0.469$, p-value = 0.000). However; using curve estimation (multiple regression), it can be seen that there is a stronger quadratic relationship between the variables. With RS L* as the independent variable, $R = 0.751$ ($R^2 = 0.564$, p-value = 0.000). With Mc L* as the independent variable, $R = 0.904$ ($R^2 = 0.816$, p-value = 0.000). This is a strong relationship and indicates that microscopic images are better at predicting the RS L* than vice versa. Figure 5.10 illustrates this relationship.

![Figure 5.10](image-url) The quadratic relationship between microscopic L* measurements and L* measurements by reflective spectrophotometry (RS). The trendline has the equation $y = 0.0188x^2 - 1.2854x + 48.505$ and $R^2 = 0.8164$. 

66
5.3.2 A Comparison of Microscopic Colour to Macroscopic Digital Image Analysis

When comparing microscopic (Mc) colour measurements to those determined by digital image analysis of macroscopic hair colour (DI) it was found that a* and b* showed no significant correlation between the two methods of measurement, while L* showed a significant, strong linear relationship between the two, with a correlation coefficient of 0.818 ($R^2 = 0.670$, p-value = 0.000). Multiple regression did not improve the prediction ability of the model.

5.3.3 A Comparison of Microscopic Colours to Reflective Spectrophotometry Clusters

Individuals were assigned to their clusters as determined in Chapter 3; LL2, LL5 and LL6. A discriminant analysis was then performed on these individuals using their microscopic colour measurement data. It was found that the percentage of cases correctly classified decreases as the number of clusters increases with LL2, LL5 and LL6 having 87.9%, 69.7% and 51.5% of cases correctly classified, respectively.

5.3.4 A Comparison of Microscopic Colours to Hair Thickness

Microscopic hair colour was compared to the estimates of hair thickness. There were significant, negative correlations between the microscopic L* values of hairs and their estimated thickness (in microns) with an $R = -0.705$ (p-value = 0.000). The a* value was positively correlated to the hair thickness ($R = 0.567$, p-value 0.000) while b* showed no significant relationship.

5.3.5 A Comparison of Macroscopic Colour to Hair Thickness

Macroscopic hair colour, measured by reflective spectrophotometry was compared to the estimates of hair thickness. Macroscopic L* and b* were found to have a moderate, negative correlation with hair thickness, with correlation coefficients of -0.648 and -0.657 (p-values = 0.000), respectively. The a* value showed a significant (p-value 0.023) but weaker correlation of -0.390.
5.3.6 Pigment Pattern Analysis

Figure 5.11 shows the results of comparing the standard deviation of the intensities of the monochrome hair images to the mean intensity value by A) image (the mean intensity of all the pixels in an image) and by B) individual (the average intensity of the nine images for each individual). The mean intensity is equivalent to the L* value for that image. There is an obvious quadratic relationship between the standard deviation and the mean intensity of the monochrome images ($R^2 = 0.741$, p-value = 0.000) which holds when individuals are examined but is not as strong ($R^2 = 0.428$), probably due to a smaller sample size. This indicates that there is less deviation in light hairs that increases as the hair gets darker until a critical point (when intensity is around 40) and then decreases again with darker hairs.
Figure 5.11 The standard deviation of the intensities of the monochrome hair images compared to the mean intensity value by individual (A) and by image (B).
5.4 Discussion

This study illustrates the relationships between macroscopic and microscopic colour, hair thickness and pigmentation pattern. It can be seen that there is a strong relationship between macroscopic and microscopic L* values that would be useful if one was trying to predict the colour of a head of hair (macroscopic) from a strand (microscopic) or if one was trying to predict hair colour phenotype by genetic analysis and either the macroscopic or microscopic hair colour was known to be more strongly associated with a genetic marker. The relationship between hair lightness (L*) and thickness is also clearly illustrated at both the macroscopic and microscopic levels.

While the reported relationships are useful and will assist in further research towards the objective measurement of hair colours, it can also been seen that there are weak or insignificant relationships between some aspects of the macroscopic and microscopic colour, particularly when examining the a* and b* colour components. These differences may be due to the fact that the number of hair strands that can be examined under the microscope will most likely be a small proportion of the head of hair and large sample sizes are difficult to obtain.

Colour measurement by digital image analysis also presents some problems. In Chapter 4, it was shown that macroscopic hair colour measured by digital images analysis does not correspond very well to the reflective spectrophotometry readings of the macroscopic hair colour and some of the same factors may be involved in measuring the colour of microscopic images; such as the ability of the camera to reproduce colour and the conversion from RGB to the L*a*b* colour space. However; the high degree of consistency in conditions under a microscope should make microscopic digital images of hairs comparable to each other and therefore more consistent in their relationships with the macroscopic colour.

The measurement of hair thickness could also be improved to increase the strength of relationships between thickness and colour. Cross-sections of the hair may be examined to determine the shape before measuring the thickness of hairs, since using the optical
depth may inadvertently use different sides of an oval-shaped hair for measurement and therefore be less precise.

The microscopic examination of the pigmentation patterns shows that hair has a smooth pigmentation distribution that becomes more variable as the hair gets darker until an intensity of about 40 and then gets smoother again (because of the saturation of pigment in the hair shaft). These results may indicate that the structure of the hair’s matrix and the distribution of pigment granules in the hair shaft (which would also be affected by the hair’s thickness) have an effect on the colour of the strands and this type of information will be important when considering which areas of the genome to examine when looking for genes that relate to hair pigmentation. For example, when studying hair colour, it may be worth looking at genes that control hair thickness since colour and thickness are correlated. Measuring additional pigmentation pattern variables, such as those considered by forensic hair examiners (Figure 2.1) may provide more information about the relationship between pigment distribution and colour.

Chapter 6 will introduce a chemical characterisation of the hair samples.
Chapter 6  Chemical Analysis of Hair Pigments

6.1  Introduction

6.1.1  Chemistry

Melanin is the biopolymer pigment that is the most directly responsible for human pigmentation [4]. While haemoglobin and dietary carotenoids contribute somewhat to the appearance of epidermal colour, they are minor constituents and have little to do with the pigmentation of hair [4].

Melanin is a polymer of quinone and indole-quinone derived units produced from the amino acid tyrosine [4, 20]. Its exact composition is unknown and the information available is inconsistent [76]. Briefly, melanin comes in several forms; two of which are relevant to human pigmentation; eumelanin and pheomelanin. Eumelanin is a black/brown pigment, while pheomelanin is a red/yellow pigment [4, 6, 51]. Studies have shown that it is both the amount of melanin present and the type of melanin that gives hair its colour [4]. Figure 6.1 shows the structure of the eumelanin and pheomelanin polymers while Figure 6.2 indicates the biosynthetic pathways of each, showing the subunits and the main enzymes involved in the synthesis of melanin, called melanogenesis.

![Figure 6.1 The structures of the eumelanin and pheomelanin polymers [76].](image-url)
Figure 6.2 The biosynthetic pathways to eumelanin and pheomelanin as established in human skin and in murine hair pigmentation [76].

As mentioned, the exact composition of the polymers is not well understood and not necessarily stable. Eumelanin is composed of 5,6-dihydroxyindole (DHI) and 5,6-dihydroxyindole-2-carboxylic acid (DHICA) derived units [76, 77]. Some hair samples [20] have been characterised as having a eumelanin composition of approximately 60% DHI-derived and 40% DHICA-derived units, however, caution is suggested in applying this result to all hair samples, given the problems in analysis of melanin and the biosynthetic mechanisms of melanin production that was discussed in Section 5.1.2. With the same caution, pheomelanin in hair is a polymer of approximately 80% 2-S-Cysteyldopa (2-S-CD) derived and 20% 5-S-Cystenyldopa (5-S-CD) derived units [20]. The cysteine residues (containing sulphur) give pheomelanin its red or yellow colour. In some animal breeds (e.g. Red Setters) pheomelanin appears red, while in others (e.g. Yellow Labrador Retrievers, certain mouse strains), pheomelanin appears yellow (See Box 6.1) [4].

Black hair has been shown to contain approximately 99% eumelanin and 1% pheomelanin, while brown and blonde hair contain approximately 95% eumelanin and
5% pheomelanin, and red hair contains 67% eumelanin and 33% pheomelanin, this unique ratio giving it its distinct, red hue [20].

The exact melanin content of blonde hair has not been examined fully and is still contentious. However the 2001 study by Borges et al. [20] indicates that blonde is a result of less overall melanin, rather than a result of a either an increased proportion of or a yellow version of pheomelanin (See Box 6.1), while red hair is a change in pigment composition. A 2005 study on the photo lightening effects of visible and ultraviolet light by Takahashi et al. [17] provides additional evidence for this. In this study, blonde hair was found to be much more sensitive to visible wavelengths of light, while red hair, known to contain a high amount of pheomelanin, was found to be more sensitive to UV light. The differential sensitivity of red and blonde to different wavelength ranges could indicate a different pigment composition.

While melanin is the pigment responsible for hair colour, it must be first synthesised by the cell and transferred to the hair. This cellular biology process was discussed in Section 5.1.2.

Box 6.1

Some animals have a yellow version of pheomelanin. The mice below have approximately the same amount of melanin but the yellow mouse is almost entirely pheomelanogenic while the brown mouse is eumelanogenic [47]. Human ‘yellow’ or blonde hair is the result of less overall melanin [20, 47].

<table>
<thead>
<tr>
<th>A/y genotype at the agouti locus</th>
<th>Non-agouti (a/a), Tyrp1b/Tyrp1b</th>
</tr>
</thead>
</table>

yellow | brown

There are several issues to be aware of when analysing or interpreting the pigment content data of hair. Melanin is an extremely difficult compound to analyse, primarily because it is a non-homogeneous biopolymer that is difficult to isolate [76, 77]. Therefore; degradation products are analysed instead, which presents new problems. The experiments by Ito and Fujita (1985) quantified pyrolle-2,3,5-tricarboxylic acid (PTCA) as the degradation product of eumelanin after a permanganate oxidation using HPLC-UV and quantified 3-amino-4-hydroxyphenylalanine (3-AHP) and 4-amino-3-hydroxyphenylalanine (4-AHP) as the degradation product of pheomelanin by HPLC-ECD (both isomers of AHP were detected, but only 3-AHP was quantified because a standard was commercially available)[77]. Factors were calculated to convert the amount of the degradation product to the amount of melanin. These conversion factors were based on the amount of PTCA and 3-AHP recovered from the degradation of sepia (cuttlefish) melanosomes (2% and 20% respectively) which gave conversion factors of 50x for PTCA and 5x for 3-AHP [77].

Later experiments in eumelanin analysis used an alkaline hydrogen peroxide degradation method that resulted in significant amounts of pyrolle-2,3-dicarboxylic acid (PDCA) being recovered and quantified, as well as PTCA [78]. These studies showed that the degradation of DHICA-derived units resulted in a 14.1% yield of PTCA and no PDCA, while the degradation of DHI-derived units resulted in a 4.82% yield of PDCA and a 0.01% yield of PTCA. Data from permanganate oxidation of eumelanin [76] also shows that PTCA primarily results from the degradation of DHICA-derived units.

Wakamatsu (2004) recommended a conversion factor of 160x for PTCA in hair due to a yield of 0.6% obtained when the concentration of PTCA in solution was measured by spectrophotometry. The low yield found in human hairs (compared to 2.2% in mice and 2.5% in sheep) indicate that most of the eumelanin in human hairs is not DHICA-derived and therefore indicates a lower activity of Tyrosinase-related protein Type 2 (TYRP2 or DCT)[76]. The pathway by which DHICA-derived units are produced by TYRP2 is illustrated in Figure 6.2. This lack of DHICA-derived eumelanin units is supported by a 2004 study by Commo et al. which found no TYRP2 protein in the
melanocytes of the human hair bulb and a barely detectable TYRP2 mRNA amplimer signal [50]. It makes sense to find little or no DHICA-derived eumelanin in hair when the enzyme that produces it (TYRP2) is not present in hair bulbs. This is in contrast to the strong presence of the enzyme and its products found in human hair follicle and skin melanocytes.

The difference in hair eumelanin compared to skin, mouse and sepia melanin presents problems for the conversion of values from PTCA yields to amounts of eumelanin and for the comparison of amounts between these tissues and species. Although the exact composition of hair melanin is unknown [79] and each method of degradation (permanganate or alkaline hydrogen peroxide) may have different yields. Until more is known about the various degradation processes, it is better to use raw PTCA values (per milligram of hair) for comparison purposes and to be conservative when comparing results between studies that use different methods.

In 2002 pheomelanin analysis, Wakamatsu et al. were able to separate the 3-amino-4-hydroxyphenylalanine isomer (3-AHP; also called 3-aminotyrosine or 3-AT) and the 4-amino-3-hydroxyphenylalanine isomer (4-AHP) and found that the 4-AHP isomer was a more specific marker for the degradation of pheomelanin [80]. Using this more specific marker, a factor of 9x is used to calculate the amount of pheomelanin from the amount of 4-AHP recovered. There are fewer reported methods for pheomelanin degradation than for eumelanin, so studies should be more comparable, however; the use of 4-AHP values, rather than pheomelanin conversions is likely to allow a more reliable comparison between studies.

A significant issue in the quantification of PTCA, PDCA and 4-AHP is the lack of a commercially available analytical standard. Only 3-AHP may be purchased and even this standard has no information about its solubility, pKa or other chemical properties. PTCA, PDCA and 4-AHP must be synthesised in-house or received from another lab. While Professor Wakamatsu (Fujita Health University, Japan) has been very generous with gifts of these standards, this lack of information, limited quantities and inconsistent availability makes optimisation of methods and new investigations difficult.
Developments in the quantification of biological pigments continue. A method for the quantification of PTCA and PDCA in skin by LC/MS/MS has been developed [81] and new degradation products for the presence of pheomelanin are being investigated [82].

6.1.3 **Aims**

The aim of this experiment was to chemically characterise the samples collected and to quantify any associations found between the amount of eumelanin and pheomelanin in hair and colour measurements taken at the macroscopic and microscopic level.

Knowing the chemical composition of the pigments in hair will assist in accurately determining of hair colour and will help to direct the decision of exactly which genes to examine in the search for associations. For example, two persons whose hair has same amount of melanin could have a different hair colour - indicating that genes outside the melanogenesis pathway may be of interest (such as hair thickness or pigment distribution).
6.2 Materials and Methods

6.2.1 Sampling

From the 202 individuals described in section 3.2.1, 58 were included in this analysis (those with hair samples available). This group includes the 36 individuals described in 5.2.1 (including three non-Europeans and two white-haired individuals) and 22 more individuals from the main sample. Approximately 30mg of hair was trimmed from each subject for chemical analysis. These samples were cut from the ends of the subjects’ hair.

6.2.2 Chemicals and Buffers Used in Eumelanin Analysis

1.05 M Sodium Hydroxide (NaOH) Ajax Finechem (Seven Hills, NSW)
30% Hydrogen Peroxide (H$_2$O$_2$) Ajax Finechem
5% Sodium Sulfite (Na$_2$SO$_3$) Sigma (Perth, WA)
6M Hydrochloric Acid (HCl) Ajax Finechem
0.01M Phosphate buffer, pH 2.1 Ortho-phosphoric acid from Ajax Finechem
100% Methanol Merck Chemicals (Kilsyth, VIC)
0.1% phthalic acid Aldrich Chemical Co. (Milwaukee, WI)

1000 ng/mL PTCA and PDCA standards (a generous gift from Professor Kazumasa Wakamatsu from the Department of Chemistry, Fujita Health University School of Health Sciences, Toyoake, Aichi, Japan)

6.2.3 Hydrogen Peroxide Degradation of Hair for Eumelanin Analysis

Sample preparation and degradation was performed according to the method described in Napolitano et al. (2000) with some differences [82]. The hair was ground in a mortar and pestle with liquid nitrogen (instead of mincing with scissors) to make handling easier and reduce loss of sample. Approximately 5mg of hair from each individual was weighed and put into a glass test tube. 950µl of 1.05M sodium hydroxide and 50µl of 30% hydrogen peroxide was added to the sample, resulting in concentrations of 1M NaOH and 1.5% H$_2$O$_2$. The test tubes were covered with Para-film and shaken for 24
hours on a Griffin Flask Shaker at room temperature. After 24 hours, 200\(\mu l\) of 5% sodium sulfite (Na\(_2\)SO\(_3\)) was added to decompose remaining H\(_2\)O\(_2\) as in Borges et al. (2001) instead of hydrogen sulfite [20]. Samples were acidified to a pH less than 4 with 143\(\mu l\) of 6M hydrochloric acid (HCl), resulting in a total volume of 1.343mL. Samples were then transferred to 1.5mL microcentrifuge tubes and centrifuged at maximum speed for 2 minutes. The supernatant was transferred to HPLC vials for analysis.

6.2.4 HPLC Conditions for Eumelanin (PTCA) Analysis

HPLC analysis was performed using Varian HPLC autosampler (Module 9100), pump (Module 9012) and UV detector (Module 9050) set to 270nm [83]. An Alltech Adsorbosphere 5\(\mu m\) 260 x 4.6 mm column (with matching guard column) was used and heated to 55\(^\circ\)C with a Biorad HPLC Column Heater. Mobile phase A was 0.01 M Phosphate Buffer (pH 2.1) and Mobile phase B was 100% methanol. Mobile phases were run in a gradient as described in [20]: A:B at 98:2 ramped evenly to 40:60 over the first 14 minutes, held at 40:60 for 6 minutes, ramped back to 98:2 over the next 5 minutes and held at 98:2 for the last 5 minutes. The total run time was 30 minutes plus a 1 minute equilibration.

6.2.5 Chemicals and Buffers Used in Pheomelanin Analysis

3-amino-4-hydroxyphenylalanine standard (3-aminotyrosine) Sigma (Perth, WA)
HPLC Grade water with 0.1% formic acid Riedel-deHaen (Germany)
20mM borate buffer (pH 9.5) Boric acid from Sigma (Perth, WA)
30mM sodium cyanide (NaCN) Ajax Finechem (Seven Hills, NSW)
20mM naphthalene-2,3-dicarboxaldehyde (NDA) Fluka (Milwaukee, WI)
Acetonitrile BDH Chemicals (Biolab, Clayton, VIC)
50mM ammonium acetate buffer (pH 6.8) Merck Chemicals (Kilsyth, VIC)

500 ng/mL each (mixed) 3-AHP and 4-AHP standards (a generous gift from Professor Kazumasa Wakamatsu from the Department of Chemistry, Fujita Health University School of Health Sciences, Toyoake, Aichi, Japan)
6.2.6 LC/MS Conditions for Pheomelanin Analysis

Small quantities of AHP are expected from the degradation of pheomelanin in hair and there is no data available on the UV absorption of AHP; therefore, more sensitive alternative detectors are employed in this analysis, such as electrochemical detection (ECD) [77]. Since ECD was not available for this experiment, an attempt was made to quantify 3-AHP with Liquid Chromatography – Mass Spectrometry (LC/MS) according to methods adapted from Nezirevic et al. (2007) [84].

The isocratic mobile phase was acetonitrile with 0.1% formic acid (pH 3.0). A ZIC-HLIC® Column (150mm x 2.1mm) was used (SeQuant, Umea, Sweden) with a Finnigan Surveyor Autosampler Plus, LC Pump Plus and LCQ Deca XP MAX module mass spectrometer (Thermo Scientific Fisher, Waltham, MA) using electrospray ionisation.

While the LC/MS method using a ZIC-HLIC® Column worked with a 3-AHP 1ppm standard, the limit of detection of this instrument (approximately 0.1ppm) was too high to be of use in the quantification of AHP from hair samples (data not shown) and was therefore, not used further.

6.2.7 HPLC Conditions for Pheomelanin using Fluorescent Detection

Methods used for the fluorescent detection of 3-AHP and 4-AHP (for the analysis of pheomelanin) were performed according to a previous study by Yang et al. (2007) [85].

First, the 3-AHP standard was put through a pre-column derivatisation [85]. 10μL of the 3-AHP standard was added into 1 mL of borate buffer (20 mM, pH9.5). After the addition of 30μL sodium cyanide (30 mM), the solutions were left to equilibrate at room temperature for 10 min. Then 10μL NDA (20 mM) solution in methanol was added and the mixture equilibrated for 30 min at 25 °C.

Chromatography was performed on the same Varian HPLC setup as described in Section 6.2.4, but with a Varian 9070 Fluorescence detector with a PMT voltage of 700.
Mobile Phase A (95%) was acetonitrile and Mobile Phase B (5%) was 50mM ammonium acetate (pH 6.8). A Supelco APS-2 Hypersil aminopropyl column was used (250mm x 4.6mm, 5µm particles) with a matching guard column. Fluorescence was detected with an excitation wavelength of 442nm and an emission wavelength of 482nm as determined by the Perkin Elmer Fluorescence Spectrophotometer LS50B.

This HPLC-FD method worked well with the 3-AHP standard and had a very low limit of detection; however the method was unable to resolve the 3-AHP and 4-AHP from a mixed standard or from actual samples. This may be due to a degradation of the standards over time (standards are not available commercially and were a gift from Professor Kazumasa Wakamatsu from the Department of Chemistry, Fujita Health University School of Health Sciences, Toyoake, Aichi, Japan) or due to the fluorescence of contaminants and undesired compounds. There is no published data on the stability of standards; therefore fresh standards and additional optimisation of this method may provide results in the future.
6.3 Results

6.3.1 Method Validation for Eumelanin (PTCA) Analysis

Standards for PTCA and PDCA (1000ng/mL) were a generous gift from Kazumasa Wakamatsu from the Department of Chemistry, Fujita Health University School of Health Sciences, Toyoake, Aichi, Japan. 1000ng/mL is equal to 1000 parts per billion (ppb) or 1 part per million (ppm). Phthalic acid was used during method optimisation since it has a retention time very near the desired compounds (Borges et al., 2001), however it was not used to quantify samples as it was found to persist through subsequent injections.

Standard curves were prepared from 50ng/mL to 1000ng/mL for PTCA and from 100ng/mL to 1000ng/mL PDCA; 50ppb and 100ppb being the lower limits of detection for PTCA and PDCA respectively. Each concentration was repeated twice and $R^2$ values for each standard curve were excellent; $R^2$ for PTCA was 0.9995 and for PDCA was 0.9951 (please see Figure 6.3 and Figure 6.4).

![Standard curve for PTCA](image)

**Figure 6.3** Standard curve for PTCA ($R^2 = 0.9995$) where $y = 34.301x - 138.04$
Retention times were approximately 9.4 minutes for PTCA and 8.3 minutes for PDCA. Degraded samples of dark and light hair were performed and analysed and typical chromatograms are shown in Figures 6.5 and 6.6.

Although PDCA was detected in the dark sample (black hair), the concentrations were so low that it was not worthwhile to continue quantifying this compound. It was not found in the fair hair at all. Blank samples showed no detectable PTCA or PDCA. Chromatograms were automatically integrated from 7 to 14 minutes.

To calculate the amount of PTCA for each milligram of hair, the following formula was used from the standard curve:

\[
\text{Concentration (ng/\mu l)} = \frac{(\text{Peak Area} + 138.04)}{34.301}
\]

\[
\text{Concentration (ng/\mu l)} \times 1.343 \text{mL} = \text{Amount of PTCA (ng)}
\]

\[
\text{Amount of PTCA (ng)/Amount of hair (mg)} = \text{ng PTCA/1 mg hair}
\]
Figure 6.5 Sample Chromatograms
A – 500 ppb PTCA Standard (9.391) and showing phthalic acid (12.748)
B – 500 ppb PDCA Standard (8.269) and showing phthalic acid (12.757)
Figure 6.6 Sample Chromatograms
C – A typical chromatogram for dark hair showing PTCA (9.391) and phthalic acid carry-over.
D – Blank sample (no hair in degradation) showing no PTCA but with phthalic acid.
For each run, a 500ppb standard was also analysed and if the calculated amount of PTCA from the standard curve was within 20% of the known amount, the run was considered valid. All runs showed a less than 4% variation for the 500ppb standard.

To validate the method with respect to variability and linearity, the following tests were performed. Tests of intra-assay variability and inter-assay variability were carried out on light and dark hair and evaluated according to the percent coefficient of variance (%CV) which is the standard deviation as a percentage of the mean. A mean of the results for two injections was used for all calculations. These results can be seen in Table 6.1.

Results for multiple injections were excellent (10% is to be expected) and overall variability for the method at 12% is acceptable and in line with the published results [20, 81]. Variation was consistent no matter if the hair was dark or light.

Table 6.1 Variability in the hair degradation and HPLC method

<table>
<thead>
<tr>
<th>Sample</th>
<th>Intra-Assay Variability (%CV)</th>
<th>Inter-Assay Variability (%CV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dark Hair</td>
<td>6.3%</td>
<td>12%</td>
</tr>
<tr>
<td>Fair Hair</td>
<td>6.5%</td>
<td>12%</td>
</tr>
</tbody>
</table>

For PTCA, a test of linearity of recovery was conducted. Various amounts of dark hair were degraded to ensure that there is a linear relationship between the amount of hair degraded and the amount of PTCA recovered. A Pearson’s test for correlation showed that the amount of hair and amount of PTCA were found to be significantly correlated (p=0.02). The variation in the final results was 9.3%CV for this test. The results are displayed in Figure 6.7.
Figure 6.7 Results of the test for linearity of PTCA recovery.

6.3.2 PTCA content of Hair Samples

The raw PTCA values and colour data for each individual can be found in Appendix 1.
6.3.3 A Comparison of PTCA Content to Colour Measured by Reflective Spectrophotometry

Figure 6.8 shows the relationship between the macroscopic L* value as measured by reflective spectrophotometry and the amount of PTCA recovered per milligram of hair. There was an expected negative correlation that indicates as hair gets lighter, there is less PTCA recovered and therefore, was less melanin present. This relationship was quite strong; however, very light hair samples (L* > 45) and white hair samples were outside the main group. Non-European samples were also outside the main group, having much more PTCA per milligram of hair.

![Figure 6.8](image_url)

**Figure 6.8** The amount of PTCA found in 1 mg of hair compared to the RS L* value. The $R^2$ for the linear relationship of the entire sample set is 0.321 and the $R^2$ for the circled group is 0.6889. (p-values = 0.000)
The strongest linear relationship between the macroscopic colour of hair, for all samples, as measured by reflective spectrophotometry and the amount of PTCA recovered was along the b* axes, as seen in Figure 6.9. No significant relationship was found between the RS a* value and the amount of PTCA recovered (R² = 0.185, p-value = 0.094).

Figure 6.9 The amount of PTCA found in 1 mg of hair compared to the RS b* value (R² = 0.6561, p-value = 0.000)
6.3.4 A Comparison of PTCA Content to the Microscopic Colour

The strongest relationship between the microscopic colour and the amount of PTCA recovered was along the L* axis. Figure 4 illustrates this negative correlation. The microscopic a* value had a moderate positive correlation with the amount of PTCA recovered ($R^2 = 0.262; p$-value = 0.001), while the microscopic b* value showed no significant correlation.

![Figure 6.10](image)

*Figure 6.10* The amount of PTCA found in 1mg of hair compared to the Mc L* value ($R^2 = 0.8769, p$-value = 0.000)
6.3.5 A Comparison of PTCA Content to Hair Thickness

A moderate significant relationship was found between the amount of PTCA recovered and the thickness of the hair as measured in Chapter 5. This relationship can be seen in Figure 6.11.

**Figure 6.11** The mean thickness of the nine hair measurements compared to the amount of PTCA recovered from 1mg of hair ($R^2 = 0.323$, p-value = 0.000)
6.3.6 PTCA Content of Reflective Spectrophotometry Clusters

Chapter 3 examined hair colour measurements and found that those measurements may be broken into objective, describable clusters. The log-likelihood method was used to break 202 European individuals into five clusters (referred to as LL5); details of these clusters can be found in Chapter 2. The 58 individuals included in this chemical analysis were also analysed according to the cluster they were assigned to in Chapter 2. The mean PTCA content (ng PTCA/mg hair) of each cluster is shown in Figure 6.12, with the standard deviation represented by the error bars. In Figure 6.12, individuals of non-European ancestry are included for comparison and are referred to as Cluster 6, although they were not part of the LL5 analysis.

![Figure 6.12](image)

**Figure 6.12** The mean content of PTCA for the LL5 reflective spectrophotometry clusters. Non-Europeans are referred to as Cluster 6 (but were not part of the LL5 analysis). The error bars represent the standard deviation in that cluster.
6.4 Discussion

The results of this experiment show how the amount of eumelanin in hair is related to the colour as measured in the CIE L*a*b* colour space by reflective spectrophotometry and microscopic analysis. As would be expected, lighter hair has less eumelanin; however, there is a stronger relationship between the amount of eumelanin present and the value of the b* (yellow) axis for macroscopic hair colour. The amount of PTCA measured is more closely related to the microscopic colour (specifically the L* axis) than to the macroscopic colour, which may be due to the different methods of colour measurement (reflective spectrophotometry compared to a digital microscopic image). Variation may also result from a chemical analysis being performed on cut ends of hair, where reflective spectrophotometry measurements were taken higher up on the head and microscopic measurements were taken along the length of the hair.

The PTCA results obtained for the clusters (Section 6.3.6) are also as expected. Clusters 5 (dark), 4 (Medium) and 2 (Fair) have progressively less of the eumelanin degradation product and all samples of European ancestry have a great deal less PTCA per milligram of hair than the non-European samples do. These results correspond reasonably well to previously published PTCA or Eumelanin results. Borges et al. (2001) also had approximately 300ng PTCA/mg hair for Black Asian hair [20], which corresponds to the non-European group in Section 6.3.6 and Napolitano et al. (2000) obtained PTCA results including the following: Black 264 ng/mg hair, Blonde 75 ng/mg hair, Light Blonde 55 ng/mg hair, Albino 10 ng/mg hair [82]. The origin of black hair in this case is unknown and the Blonde and Light Blonde are higher than the results obtained in this study, which may be a result of variation in methods, instrumentation or in the assignment of colour categories. It is interesting to note that the results for Type I Albino hair (Tyrosinase-negative) in Napolitano et al. (2000) were almost the same as those for White hair in this study – approximately 10ng PTCA per milligram of hair.

While the relationship between colour and eumelanin content of the hair is obvious, it is not perfect. Some variation in quantification may be responsible, but it may also be the case that there are other contributors to hair colour, such as the structure and thickness of the keratin matrix of the hair shaft, the distribution of the eumelanin pigment in the
hair and the amount and ratio of the pheomelanin pigment. Although the pheomelanin component of these samples was not quantified, the thickness of the hair was measured and was found to be moderately related to the amount of PTCA recovered. Two hairs with the same amount of eumelanin per milligram of hair may appear to be a different colour if the hair is thinner and the pigment is, therefore, more densely packed or is distributed in a different way.

While it is unfortunate that the quantification of pheomelanin content of these samples was unsuccessful and further work in this area would be beneficial, it should be noted that this study is primarily concerned with the characterisation of hair along the dark-fair continuum, where pheomelanin is a very minor component (approximately 5% of the pigment content [20]) and the ratio between eumelanin and pheomelanin less important than if one were studying red hair.
Chapter 7  Genetic Analysis

7.1  Introduction

7.1.1  Genetic Control of Pigmentation

It has been suggested that most pigmentation variation is probably the result of a few highly penetrant alleles [10], however, the search for these alleles and the genes they are hidden in is far from over. From studies of epidermal and iris pigmentation, as well as from murine models, we know some of the genes involved, or likely to be involved, in human pigmentation. Protein-coding genes TYR [86], TYRP 1 and 2 [86, 87], the P gene (OCA2) [86, 88, 89], MC1R [4, 14, 18, 86, 90], ASIP [40, 44], MATP [45], POMC [91], SILV [51, 86], MYO5A and RAB27A [5, 86], as well as transcription factors such as MITF or SOX10 have been implicated in human pigmentation [3, 6, 51].

Nonsynonymous mutations may lead to peptides with greater or lesser activity. Tyrosinase, the rate-limiting enzyme in melanin synthesis, has been found to have at least ten times greater activity in melanocytes derived from ‘black’ skin compared to those derived from ‘white’ skin [86], while the amount of protein remained the same. Normal variation of human pigmentation may also be the result of differences in regulatory elements at the transcriptional, translational or posttranslational levels [40], rather than in coding differences. For example, a SNP in the 3’ untranslated region of ASIP is correlated with darker pigmentation [44]. So far, mutations in the melanocortin-1 receptor have been the only variations directly correlated with hair colour differences; specifically red hair [4, 6, 18, 86, 90].

The following tables (Table 7.1 to Table 7.4) are adapted from the Albinism Database [92] and other relevant studies (cited in square brackets in the table) and indicate genes thought to be most involved in normal human pigmentation variation [92, 93] due to their disease-state (the appearance of individuals with a defective gene). Also listed are the chromosomal locations of the gene, the human disease or phenotype associated with a non-functioning protein, or a protein with reduced function, and the function or

95
potential function of the protein in normal pigmentation. There are likely to be other genes involved in different aspects of pigmentation (candidate pigmentation genes), such as MLPH [94] or SLC24A5 [95]; genes that are logically connected to the pigmentation pathway but do not have phenotypes as obvious as those in Tables 7.1 to 7.4. Several of the genes tested in this study are of this type.

Table 7.1 Genes involved in the synthesis of melanin.

<table>
<thead>
<tr>
<th>Human Locus</th>
<th>Alternate Name</th>
<th>Human Chromosome</th>
<th>Associated Disease</th>
<th>Protein Encoded</th>
<th>Function in Pigmentation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TYR</strong> [86]</td>
<td></td>
<td>11q21</td>
<td>oculocutaneous albinism type 1 (OCA1)</td>
<td>tyrosinase</td>
<td>Oxidises tyrosine and dopa to dopaquinone and polymerises DHI units into eumelanin</td>
</tr>
<tr>
<td><strong>TYRP1</strong> [86, 87]</td>
<td></td>
<td>9p23</td>
<td>oculocutaneous albinism type 3 (OCA3) Rufous albinism</td>
<td>tyrosinase related protein 1 (TRP1)</td>
<td>Part of the tyrosinase complex (stabilising factor) and DHICA oxidase; incorporating DHICA units into eumelanin</td>
</tr>
<tr>
<td><strong>TYRP2</strong> [86, 87]</td>
<td><strong>DCT</strong></td>
<td>13q31-q32</td>
<td>Unknown</td>
<td>tyrosinase related protein 2 (TRP2)</td>
<td>DOPAchrome tautomerase – modifying dopachrome to DHICA units</td>
</tr>
<tr>
<td><strong>MATP</strong> [45]</td>
<td></td>
<td>5p</td>
<td>oculocutaneous albinism type 4 (OCA4)</td>
<td>Membrane-associated transporter protein</td>
<td></td>
</tr>
<tr>
<td><strong>Pmel-17</strong> [51, 86]</td>
<td><strong>SILV, gp100</strong></td>
<td>12q13-q14</td>
<td>Unknown</td>
<td>gp100/silver protein, Pmel17</td>
<td>melanosomal matrix protein – specific marker for melanocytes</td>
</tr>
</tbody>
</table>

Table 7.2 Genes involved in melanosome structure and transport.

<table>
<thead>
<tr>
<th>Human Locus</th>
<th>Alternate Name</th>
<th>Human Chromosome</th>
<th>Associated Disease</th>
<th>Protein Encoded</th>
<th>Function in Pigmentation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MYO5A</strong> [5, 86]</td>
<td></td>
<td>15q21</td>
<td>Griscelli syndrome</td>
<td>myosin type Va</td>
<td>Movement of the melanosome</td>
</tr>
<tr>
<td><strong>RAB27A</strong> [5, 86]</td>
<td></td>
<td>15q21</td>
<td>Griscelli syndrome</td>
<td>RAS associated protein</td>
<td>Movement of the melanosome</td>
</tr>
<tr>
<td><strong>P</strong> [86, 88, 89]</td>
<td><strong>OCA2</strong></td>
<td>15q11.2-q12</td>
<td>oculocutaneous albinism type 2 (OCA2)</td>
<td>Unknown; on melanosome membrane</td>
<td>Many proposed; ion exchange, pH control. Main gene involved in iris pigmentation</td>
</tr>
</tbody>
</table>
Table 7.3 Genes involved in the type of melanin produced (Eumelanin/Pheomelanin switch).

<table>
<thead>
<tr>
<th>Human Locus</th>
<th>Human Chromosome</th>
<th>Associated Phenotype</th>
<th>Protein Encoded</th>
<th>Function in Pigmentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>MC1R [4, 14, 86, 90]</td>
<td>16q24.3</td>
<td>hair colour/skin type</td>
<td>Melanocortin 1 receptor</td>
<td>Eumelanin / pheomelanin switch</td>
</tr>
<tr>
<td>ASIP [40, 44]</td>
<td>20q11.2</td>
<td>hair colour/skin type</td>
<td>agouti signal protein (ASIP)</td>
<td>Eumelanin / pheomelanin switch</td>
</tr>
<tr>
<td>POMC [91]</td>
<td>2p23.3</td>
<td>Red Hair</td>
<td>Proopiomelanocortin; including Melanocyte Stimulating Hormone (MSH) Ligand to MCIR</td>
<td>Eumelanin / pheomelanin switch (and endocrine functions)</td>
</tr>
</tbody>
</table>

Table 7.4 Transcription factors that may be involved in human pigmentation.

<table>
<thead>
<tr>
<th>Human Locus</th>
<th>Human Chromosome</th>
<th>Associated Phenotype</th>
<th>Protein Encoded</th>
<th>Function in Pigmentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>MITF [3, 6, 51]</td>
<td>3p12-14</td>
<td>Waardenburg syndrome type 2</td>
<td>transcription factor</td>
<td>melanocyte differentiation</td>
</tr>
<tr>
<td>SOX10 [3, 6, 51]</td>
<td>22q13</td>
<td>Waardenburg-Shah Syndrome</td>
<td>SRY-box containing gene 10</td>
<td>transcriptional activator</td>
</tr>
</tbody>
</table>

Researchers are beginning to realise that there are fundamental differences between the genetic control of mouse and human pigmentation and between the pigmentation of human epidermis, iris and hair. It has recently been shown that Tyrosinase-Related Protein Type 2 (TYRP 2) is not expressed in hair follicle melanocytes found in the hair bulb, the site of melanogenesis for hair pigmentation [50]. Another study [91] has shown that there are major differences in the expression of the POMC/MC1R signalling pathway in hair follicle melanocytes, compared to epidermal melanocytes. In addition, several studies have shown that MC1R interacts with other genes, especially the P-gene [7, 15, 96] and possibly ASIP [40] and is not solely responsible for all hair colour variation [4, 40]. Much work needs to be done to elucidate the genetic contributors to normal hair colour variation.
7.1.2 Genetic Association Studies of Hair Colour

Sections 2.3.2 and 2.3.3 have discussed the results of and issues with genetic studies of hair colour. Like this investigation, these studies are typically candidate-gene association studies; where single nucleotide polymorphisms are examined in genes that have a logical connection to the pigmentation pathway and SNPs found to have significant associations may be mutations that cause variation in the resulting pigmentation, or they may be in linkage disequilibrium with mutations that do.

Population structure can lead to false positive associations when particular SNPs are ancestry-informative rather than phenotype specific [9]. For example, a SNP that has nothing to do with pigmentation may be associated with African ancestry and if there are persons of African ancestry in the sample, that SNP may end up being positively associated with darker hair, since people with African ancestry often have darker hair.

Choosing individuals with European ancestry limits the possibility of population structure interfering in the results of the genetic analysis.

7.1.3 Aims

It is hypothesised that there are genetic variants specifically associated with an objectively defined Blonde phenotype. Chapter 3 has illustrated one way of grouping hair colours into “clusters” according to their colour as measured by reflective spectrophotometry.

A total of 58 Single Nucleotide Polymorphisms (SNPs) were genotyped on 190 individuals of European ancestry and this Chapter will report on preliminary associations found with the clusters determined in Chapter 3 as well as quantitative measurements of CIE L*, a* and b*. It is recognised that this is a small sample, but it is well-characterised with objective measurements and will provide an indication of which genes and polymorphism should be further considered. While there are many additional tests that could be conducted, the following are focused on associations with fair/blonde hair in order to best achieve the study aims.
7.2 Materials and Methods

7.2.1 Sampling

The genetic analysis included 190 individuals of European ancestry that volunteered a DNA sample, from the group described in 3.2.1. The two white-haired individuals were not included. DNA samples were collected via buccal swab (four per individual) using either Omniswabs (Fitzco, Spring Park, MN) or cotton swabs (Copan Italia, Brescia, Italy). Omniswab heads were ejected into microcentrifuge tubes upon collection, allowed to dry and the tubes shut. Cotton swabs were placed in their sheaths with the ends cut off to allow drying and put in a sealed envelope.

7.2.2 Single Nucleotide Polymorphisms

The SNPs tested in this analysis are listed in Table 7.5 according to the SNP_ID number that can be used to find information about each polymorphism in the Entrez SNP database [97]. SNPs were chosen by Angela van Daal’s research group at Bond University (Queensland, Australia) for a previous study of pigmentation associations and also covered most areas of interest for this research [98]. These SNPs were selected to cover candidate-pigmentation genes [99]. Candidate-pigmentation genes are those that can be connected to the melanin production pathway, such as those in the melanin synthesis pathway (e.g. TYR, MATP) or genes involved in melanosome structure (e.g. OCA2).
Table 7.5 Single Nucleotide Polymorphisms tested for association with hair colour and the genes in which they are found.

<table>
<thead>
<tr>
<th>SNP_ID</th>
<th>Gene</th>
<th>SNP_ID</th>
<th>Gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>RS6453373</td>
<td>AP3B1</td>
<td>RS1805007</td>
<td>MC1R</td>
</tr>
<tr>
<td>RS1801244</td>
<td>ATP7B</td>
<td>RS885479</td>
<td>MC1R</td>
</tr>
<tr>
<td>RS1801243</td>
<td>ATP7B</td>
<td>RS1805005</td>
<td>MC1R</td>
</tr>
<tr>
<td>RS7334118</td>
<td>ATP7B</td>
<td>RS2228479</td>
<td>MC1R</td>
</tr>
<tr>
<td>RS1801249</td>
<td>ATP7B</td>
<td>RS4468625</td>
<td>MGRN1</td>
</tr>
<tr>
<td>RS732774</td>
<td>ATP7B</td>
<td>RS3751109</td>
<td>MLPH</td>
</tr>
<tr>
<td>RS17435026</td>
<td>ATP7B</td>
<td>RS3817362</td>
<td>MLPH</td>
</tr>
<tr>
<td>RS6107308</td>
<td>ATRN</td>
<td>RS2292884</td>
<td>MLPH</td>
</tr>
<tr>
<td>RS3886999</td>
<td>ATRN</td>
<td>RS3751107</td>
<td>MLPH</td>
</tr>
<tr>
<td>RS7782078</td>
<td>ATRN</td>
<td>RS11883500</td>
<td>MLPH</td>
</tr>
<tr>
<td>RS16876571</td>
<td>DTNB1</td>
<td>RS6435927</td>
<td>MREG</td>
</tr>
<tr>
<td>RS17470454</td>
<td>DTNB1</td>
<td>RS1058219</td>
<td>MYO5A</td>
</tr>
<tr>
<td>RS17336807</td>
<td>Egfr</td>
<td>RS16964944</td>
<td>MYO5A</td>
</tr>
<tr>
<td>RS11543848</td>
<td>Egfr</td>
<td>RS1724577</td>
<td>MYO5A</td>
</tr>
<tr>
<td>RS2296434</td>
<td>HPS1</td>
<td>RS1052030</td>
<td>MYO7A</td>
</tr>
<tr>
<td>RS2296436</td>
<td>HPS1</td>
<td>RS2276288</td>
<td>MYO7A</td>
</tr>
<tr>
<td>RS3747129</td>
<td>HPS4</td>
<td>RS7117511</td>
<td>MYO7A</td>
</tr>
<tr>
<td>RS2014410</td>
<td>HPS4</td>
<td>RS2305253</td>
<td>OCA2</td>
</tr>
<tr>
<td>RS1894706</td>
<td>HPS4</td>
<td>RS1800414</td>
<td>P(OCA2)</td>
</tr>
<tr>
<td>RS1894704</td>
<td>HPS4</td>
<td>RS28945095</td>
<td>PAX3</td>
</tr>
<tr>
<td>RS752330</td>
<td>HPS4</td>
<td>RS729421</td>
<td>RABGTA</td>
</tr>
<tr>
<td>RS7128017</td>
<td>HPS5</td>
<td>RS17118154</td>
<td>SILV (Pmel-17)</td>
</tr>
<tr>
<td>RS14024</td>
<td>KRT1</td>
<td>RS1426654</td>
<td>SLC24A5</td>
</tr>
<tr>
<td>RS2634041</td>
<td>KRT2</td>
<td>RS26722</td>
<td>SLC45A2 (MATP)</td>
</tr>
<tr>
<td>RS638043</td>
<td>KRT2</td>
<td>RS16891982</td>
<td>SLC45A2 (MATP)</td>
</tr>
<tr>
<td>RS7541041</td>
<td>LYST</td>
<td>RS1800422</td>
<td>TYR</td>
</tr>
<tr>
<td>RS3768067</td>
<td>LYST</td>
<td>RS1042602</td>
<td>TYR</td>
</tr>
<tr>
<td>RS10465613</td>
<td>LYST</td>
<td>RS16929374</td>
<td>TYRP1</td>
</tr>
<tr>
<td>RS7524261</td>
<td>LYST</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
7.2.3 **DNA Extraction, Quantification, Preparation and Genotyping**

DNA was extracted from two buccal swabs from each individual using the Qiagen Gentra Puregene Buccal Cell Kit according to the manufacturer’s instructions with some modification. The two swab heads were placed into one microcentrifuge tube (cotton swabs had the heads cut off the wooden shaft). 600 μL of Qiagen cell lysis solution was used with cotton swabs and 800 μL with Omniswabs to completely cover both swab heads. 3 μL of Proteinase K (>600 mAU/mL; Qiagen) and RNase A (17500U/mL; Qiagen) was used with the cotton swabs and 4 μL with the Omniswabs. One mAU is the activity that releases folin-positive amino acids and peptides corresponding to 1 µmol tyrosine per minute \[100\]. 200 μL of Protein Precipitation solution were used for both swab types. 600 μL of isopropanol was used to precipitate DNA according to the kit procedure and samples were hydrated in 40 μL of sterile, distilled and deionised water.

DNA samples were quantified using the Quantifiler Human DNA Quantification kit (Applied Biosystems Australia, Victoria, Australia), a real-time qPCR method which was performed according to standard protocols at the Victoria Police Forensic Services Department \[101, 102\].

If samples had more than 10 ng/μL of DNA, they were diluted to this concentration using sterile ddH2O. If the concentration was between 5 ng/μL and 10 ng/μL, the samples were concentrated using an isopropanol DNA precipitation \[103\] as follows; 1 Volume of DNA sample was added to 0.1 Volume of Sodium Acetate and 1 Volume of isopropanol and held at -20°C overnight. The sample was centrifuged at 14 000g and the supernatant removed. The precipitated DNA pellet was washed with 70% ethanol and dried. The sample was then re-hydrated in the appropriate amount of sterile ddH2O to make 10 ng/μL (assuming 100% recovery). If samples had less than 5 ng/μL of DNA, the last two swabs were extracted and the procedure conducted again. The above procedures resulted in successful DNA recovery for each individual.

10 μL of each DNA sample with a concentration of 10 ng/μL were loaded into 96-well plates and were sent to the Australian Genome Research Facility for genotyping via the
Sequenom MassARRAY system. A total of 58 SNPs were genotyped, of which, 46 were found to be polymorphic in this population.

7.2.4 Software and Statistical Analysis

Genetic analysis was performed with software by Golden Helix (Bozeman, MT); HelixTree and the SNP Variation Suite (free trial). Allele frequency data was analysed using Chi squared tests ($\chi^2$) and genotype data was analysed using Fisher’s Exact Test. Hardy-Weinberg Equilibrium and linkage disequilibrium were also assessed using this software. Regression analysis to test for second-order interaction terms was also performed with the HelixTree software.

No corrections were made for multiple analyses and the results, therefore, may reflect some degree of false positive associations. Bonferroni adjustments eliminated any significant associations, but this method is thought to be overly conservative for this study [104, 105]. Previous hair colour genetics studies do not mention the use of multiple analysis correction [38, 45].

Due to the small sample size in this study, $\alpha = 0.1$ was used as a relaxed requirement of significance.
7.3 Results

7.3.1 Polymorphic Single Nucleotide Polymorphisms

Of the 58 SNPs tested, 46 were found to be polymorphic. Two SNPs, rs17336807 and rs11543848 are recognised as the same polymorphism which is now referred to as rs2227983 [106]. Eleven SNPs were found to be non-polymorphic. All of these were expected and have been reported as being fixed in European populations by the International HapMap project [107]. However, one SNP that has been reported as fixed in Europeans by HapMap was found to be polymorphic in this sample. The SNP rs1426654 had a minor allele frequency of 0.014 in this sample, which is consistent with other research [95].

Table 7.6 shows the minor allele frequency of the typed SNPs as well as the call (or success) rate of genotyping. Raw genotype data may be found in Appendix 5.

**Table 7.6**: SNP results showing the minor allele frequency and the call rate – non-polymorphic SNPs have a minor allele frequency of 0.00.

<table>
<thead>
<tr>
<th>Marker</th>
<th>Minor Allele Freq.</th>
<th>Call Rate</th>
<th>Marker</th>
<th>Minor Allele Freq.</th>
<th>Call Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>RS1042602</td>
<td>0.339</td>
<td>0.979</td>
<td>RS2228479</td>
<td>0.067</td>
<td>0.984</td>
</tr>
<tr>
<td>RS10465613</td>
<td>0.000</td>
<td>0.995</td>
<td>RS2276288</td>
<td>0.426</td>
<td>0.989</td>
</tr>
<tr>
<td>RS1052030</td>
<td>0.353</td>
<td>0.984</td>
<td>RS2292884</td>
<td>0.213</td>
<td>0.989</td>
</tr>
<tr>
<td>RS1058219</td>
<td>0.169</td>
<td>0.979</td>
<td>RS2296434</td>
<td>0.080</td>
<td>0.984</td>
</tr>
<tr>
<td>RS1061472</td>
<td>0.428</td>
<td>0.989</td>
<td>RS2296436</td>
<td>0.080</td>
<td>0.984</td>
</tr>
<tr>
<td>RS11543848</td>
<td>0.293</td>
<td>0.968</td>
<td>RS2305253</td>
<td>0.000</td>
<td>0.995</td>
</tr>
<tr>
<td>RS11883500</td>
<td>0.147</td>
<td>0.968</td>
<td>RS2634041</td>
<td>0.000</td>
<td>0.995</td>
</tr>
<tr>
<td>RS14024</td>
<td>0.297</td>
<td>0.984</td>
<td>RS26722</td>
<td>0.021</td>
<td>0.984</td>
</tr>
<tr>
<td>RS1426654</td>
<td>0.014</td>
<td>0.968</td>
<td>RS28945095</td>
<td>0.005</td>
<td>0.984</td>
</tr>
<tr>
<td>RS16876571</td>
<td>0.013</td>
<td>0.995</td>
<td>RS3747129</td>
<td>0.144</td>
<td>0.984</td>
</tr>
<tr>
<td>RS16891982</td>
<td>0.051</td>
<td>0.989</td>
<td>RS3751107</td>
<td>0.141</td>
<td>0.968</td>
</tr>
<tr>
<td>RS16929374</td>
<td>0.000</td>
<td>0.995</td>
<td>RS3751109</td>
<td>0.144</td>
<td>0.989</td>
</tr>
<tr>
<td>RS16964944</td>
<td>0.000</td>
<td>0.995</td>
<td>RS3768067</td>
<td>0.000</td>
<td>0.995</td>
</tr>
<tr>
<td>RS17118154</td>
<td>0.003</td>
<td>0.984</td>
<td>RS3817362</td>
<td>0.091</td>
<td>0.863</td>
</tr>
<tr>
<td>RS1724577</td>
<td>0.000</td>
<td>0.995</td>
<td>RS3886999</td>
<td>0.080</td>
<td>0.989</td>
</tr>
<tr>
<td>RS17336807</td>
<td>0.293</td>
<td>0.989</td>
<td>RS4468625</td>
<td>0.000</td>
<td>0.995</td>
</tr>
<tr>
<td>RS17435026</td>
<td>0.429</td>
<td>0.958</td>
<td>RS5752330</td>
<td>0.130</td>
<td>0.989</td>
</tr>
<tr>
<td>RS17470454</td>
<td>0.059</td>
<td>0.989</td>
<td>RS6107308</td>
<td>0.005</td>
<td>0.989</td>
</tr>
<tr>
<td>RS17782078</td>
<td>0.080</td>
<td>0.989</td>
<td>RS638043</td>
<td>0.147</td>
<td>0.968</td>
</tr>
<tr>
<td>RS1800414</td>
<td>0.000</td>
<td>0.995</td>
<td>RS6435927</td>
<td>0.212</td>
<td>0.979</td>
</tr>
<tr>
<td>RS1800422</td>
<td>0.340</td>
<td>0.984</td>
<td>RS6453373</td>
<td>0.067</td>
<td>0.984</td>
</tr>
<tr>
<td>RS1801243</td>
<td>0.487</td>
<td>0.984</td>
<td>RS7117511</td>
<td>0.003</td>
<td>0.989</td>
</tr>
<tr>
<td>RS1801244</td>
<td>0.484</td>
<td>0.984</td>
<td>RS7128017</td>
<td>0.120</td>
<td>0.968</td>
</tr>
<tr>
<td>RS1801249</td>
<td>0.420</td>
<td>0.989</td>
<td>RS729421</td>
<td>0.385</td>
<td>0.984</td>
</tr>
<tr>
<td>RS1805005</td>
<td>0.133</td>
<td>0.968</td>
<td>RS732774</td>
<td>0.420</td>
<td>0.989</td>
</tr>
<tr>
<td>RS1805007</td>
<td>0.110</td>
<td>0.984</td>
<td>RS7334118</td>
<td>0.016</td>
<td>0.984</td>
</tr>
<tr>
<td>RS1894704</td>
<td>0.090</td>
<td>0.989</td>
<td>RS7524261</td>
<td>0.000</td>
<td>0.995</td>
</tr>
<tr>
<td>RS1894706</td>
<td>0.090</td>
<td>0.989</td>
<td>RS7541041</td>
<td>0.000</td>
<td>0.995</td>
</tr>
<tr>
<td>RS2014410</td>
<td>0.398</td>
<td>0.984</td>
<td>RS885479</td>
<td>0.055</td>
<td>0.963</td>
</tr>
</tbody>
</table>
7.3.2 **Hardy-Weinberg Equilibrium and Linkage Disequilibrium**

All SNPs except three were in Hardy-Weinberg Equilibrium, which may be due to the effects of natural selection or non-random mating. SNPs not in HWE were rs6453373, rs16891982 and rs17435026.

Figure 7.1 illustrates the low amount of linkage disequilibrium found amongst the markers used in this study (blue means a low amount of LD). The plot takes into account which markers are on the same chromosome and which may be linked.
Figure 7.1 Linkage disequilibrium plot of SNP markers that have been mapped according to their chromosomal locations.
7.3.3 Tests for Associations with Categorical Variables (Clusters)

Several genetic models were tested for SNPs that were polymorphic in this population. Individuals in the sample were treated as cases or controls, with cases being the particular hair colour phenotype under study and controls being the rest of the population. The Odds Ratio (OR) is reported for the effect that the minor allele or particular genotype being considered has on the case phenotype. An odds ratio of 1 means that this allele or genotype is equally likely to be found in both cases and controls. An odds ratio greater than one means that this particular allele or genotype is more likely to be found in the cases than in the controls. An odds ratio less than one means that this particular allele or genotype is more likely to be found in the controls than in the cases (this is the case when the major allele is associated with the phenotype being considered).

The SNP being considered, the minor allele, the odds ratio, the gene that the SNP can be found in and the amino acid substitution it causes are reported for each of the models tested (Table 7.7 Table 7.15). In Tables 7.6 to 7.14; N equals the number of cases, D represents the minor allele and d represents the major allele.

The whole population was considered first. It is well known that variations in the gene MC1R contribute to Red hair colour [18], so the first allelic test used individuals in Cluster 1 (Red) as cases (Table 7.7). Table 7.8 shows the SNPs that were found to be significantly associated with Cluster 2 (Blonde/Fair) at the allelic level.

**Table 7.7** SNPs found to be significantly associated with LL5 Red (Cluster 1) at the $\alpha = 0.1$ level for the allelic model (N = 8)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1805007</td>
<td>T</td>
<td>13.1</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs1800422</td>
<td>A</td>
<td>3.4</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
</tbody>
</table>
Table 7.8 SNPs found to be significantly associated with LL5 Fair/Blonde (Cluster 2) at the $\alpha = 0.1$ level for the allelic model (N = 32)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs16876571</td>
<td>A</td>
<td>7.7</td>
<td>DTNBP1</td>
<td>H297Y</td>
</tr>
<tr>
<td>rs2276288</td>
<td>A</td>
<td>1.6</td>
<td>MYO7A</td>
<td>C1628S</td>
</tr>
</tbody>
</table>

The following tests use Cluster 2 (Fair/Blonde) as the cases for a dominant model of association (Table 7.9) and a recessive model (Table 7.10).

Table 7.9 SNPs found to be significantly associated with LL5 Fair/Blonde (Cluster 2) at the $\alpha = 0.1$ level for the dominant model (N = 32)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs16876571</td>
<td>A</td>
<td>8.0</td>
<td>DTNBP1</td>
<td>H297Y</td>
</tr>
<tr>
<td>rs2276288</td>
<td>A</td>
<td>2.4</td>
<td>MYO7A</td>
<td>C1628S</td>
</tr>
<tr>
<td>rs1805007</td>
<td>T</td>
<td>2.1</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
</tbody>
</table>

Table 7.10 SNPs found to be significantly associated with LL5 Fair/Blonde (Cluster 2) at the $\alpha = 0.1$ level for the recessive model (N = 32)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1800422</td>
<td>A</td>
<td>3.1</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs17435026</td>
<td>A</td>
<td>2.1</td>
<td>ATP7B</td>
<td>R832K</td>
</tr>
</tbody>
</table>
Since the variations in MC1R leading to red hair are well known and can already be tested for [14, 18, 37-41, 108], additional tests were conducted with the eight individuals from Cluster 1 (Red) removed from the population. This was done to independently test for associations with Fair/Blonde hair colour at the allelic level (Table 7.11) and with a dominant (Table 7.12) or recessive model (Table 7.13).

**Table 7.11** SNPs found to be significantly associated with LL5 Fair/Blonde (Cluster 2) at the \( \alpha = 0.1 \) level for the allelic model with Cluster 1 removed (\( N = 32 \))

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs16876571</td>
<td>A</td>
<td>7.3</td>
<td>DTNBP1</td>
<td>H297Y</td>
</tr>
<tr>
<td>rs1805007</td>
<td>T</td>
<td>2.3</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs2276288</td>
<td>A</td>
<td>1.7</td>
<td>MYO7A</td>
<td>C1628S</td>
</tr>
<tr>
<td>rs1800422</td>
<td>A</td>
<td>1.7</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
</tbody>
</table>

**Table 7.12** SNPs found to be significantly associated with LL5 Fair/Blonde (Cluster 2) at the \( \alpha = 0.1 \) level for the dominant model with Cluster 1 removed (\( N = 32 \))

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs16876571</td>
<td>A</td>
<td>7.6</td>
<td>DTNBP1</td>
<td>H297Y</td>
</tr>
<tr>
<td>rs1805007</td>
<td>T</td>
<td>2.6</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs2276288</td>
<td>A</td>
<td>2.4</td>
<td>MYO7A</td>
<td>C1628S</td>
</tr>
</tbody>
</table>

**Table 7.13** SNPs found to be significantly associated with LL5 Fair/Blonde (Cluster 2) at the \( \alpha = 0.1 \) level for the recessive model with Cluster 1 removed (\( N = 32 \))

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1800422</td>
<td>A</td>
<td>3.8</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
</tbody>
</table>
Tests for genetic associations with LL5 Cluster 5 (dark hair) were conducted to see if any SNPs had the effect of contributing to darker hair colours and may indicate an opposing process than that which contributes to light hair colours. Table 7.14 and Table 7.15 show the associations found in an allelic and a dominant model. Three SNPs showed associations with LL5 Cluster 5 (Dark) and all were different SNPs than had been associated with Red or Blonde/Fair hair colour (and not simply the opposite allele). However, two SNPs were in ATP7B, a gene that had previously shown a different SNP to be associated with Blonde/Fair hair (See Table 7.10).

**Table 7.14** SNPs found to be significantly associated with LL5 Dark (Cluster 5) at the $\alpha = 0.1$ level for the allelic model ($N = 86$)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs16891982</td>
<td>C</td>
<td>3.86</td>
<td>MATP</td>
<td>L374F</td>
</tr>
</tbody>
</table>

**Table 7.15** SNPs found to be significantly associated with LL5 Dark (Cluster 5) at the $\alpha = 0.1$ level for the dominant model ($N = 86$)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Minor Allele</th>
<th>OR</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs16891982</td>
<td>C</td>
<td>3.46</td>
<td>MATP</td>
<td>L374F</td>
</tr>
<tr>
<td>rs1801243</td>
<td>G</td>
<td>0.49</td>
<td>ATP7B</td>
<td>S406A</td>
</tr>
<tr>
<td>rs1801244</td>
<td>C</td>
<td>0.49</td>
<td>ATP7B</td>
<td>V456L</td>
</tr>
</tbody>
</table>


**7.3.4 Tests for Associations with Quantitative Variables**

Associations with the quantitative variables L, a* and b* were examined next. The mean of 15 reflective spectrophotometry measurements (as described in Section 3.2.2) was used to test for associations. A significance level of $\alpha = 0.1$ was used. The Corr/Trend (Correlation/Trend) is a linear regression test that will show the $p$-value for the dependent variable (L*, a* or b*) value having any correlation with or “trend” which depends upon the count value of the genotype. The Corr/Trend R value indicates the degree and direction of this trend. The F-test is also used, as this test is on whether the distributions of the dependent variable (L*, a* or b*) within each category are significantly different between the various categories of the predictor variable (genotypes DD, Dd and dd). These tests were performed on the whole sample; no individuals (like the Reds in 7.3.2) were removed.

Table 7.16 illustrates allelic associations with L*. It shows that five SNPs have a positive correlation with L*, meaning the minor allele is associated with lighter hair (higher L* values) while rs16891982 shows a negative correlation with L* (the minor allele is associated with darker hair).

**Table 7.16 SNPs that have significant allelic associations with L* (D vs. d)**

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs16891982</td>
<td>0.005</td>
<td>-0.144</td>
<td>0.005</td>
<td>MATP</td>
<td>L374F</td>
</tr>
<tr>
<td>rs1800422</td>
<td>0.005</td>
<td>0.144</td>
<td>0.005</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs1805007</td>
<td>0.029</td>
<td>0.113</td>
<td>0.029</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs16876571</td>
<td>0.040</td>
<td>0.106</td>
<td>0.040</td>
<td>DTNBP1</td>
<td>H297Y</td>
</tr>
<tr>
<td>rs17782078</td>
<td>0.079</td>
<td>0.091</td>
<td>0.079</td>
<td>ATRN</td>
<td>I426T</td>
</tr>
<tr>
<td>rs3886999</td>
<td>0.079</td>
<td>0.091</td>
<td>0.079</td>
<td>ATRN</td>
<td>R1152K</td>
</tr>
</tbody>
</table>
The F-test statistic shown in Table 7.17 is used to test whether the distribution of the dependent variable (L*) within each category are significantly different between the various possible genotypes (DD, Dd or dd). The SNPs shown in Table 7.17 have significant differences of L* between the three possible genotypes.

**Table 7.17** A test for genotypic associations with L* (DD vs. Dd vs. dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1800422</td>
<td>0.007</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs16891982</td>
<td>0.035</td>
<td>MATP</td>
<td>L374F</td>
</tr>
<tr>
<td>rs16876571</td>
<td>0.039</td>
<td>DTNBP1</td>
<td>H297Y</td>
</tr>
<tr>
<td>rs3751107</td>
<td>0.041</td>
<td>MLPH</td>
<td>G172D</td>
</tr>
<tr>
<td>rs3751109</td>
<td>0.056</td>
<td>MLPH</td>
<td>L153P</td>
</tr>
<tr>
<td>rs11883500</td>
<td>0.062</td>
<td>MLPH</td>
<td>T289I</td>
</tr>
<tr>
<td>rs1052030</td>
<td>0.068</td>
<td>MYO7A</td>
<td>T16S</td>
</tr>
</tbody>
</table>

Table 7.18 and Table 7.19 show significant associations between various SNPs and L* in a dominant and recessive model, respectively. As mentioned above, positive correlations (R) indicate that the minor allele is associated with higher (lighter) L* values while negative correlations (R) indicate that the minor allele is associated with lower (darker) L* values.

**Table 7.18** SNPs that have a significant association with L* in dominant model (DD, Dd vs dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs16891982</td>
<td>0.012</td>
<td>-0.184</td>
<td>0.012</td>
<td>MATP</td>
<td>L374F</td>
</tr>
<tr>
<td>rs1805007</td>
<td>0.036</td>
<td>0.153</td>
<td>0.036</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs16876571</td>
<td>0.039</td>
<td>0.150</td>
<td>0.039</td>
<td>DTNBP1</td>
<td>H297Y</td>
</tr>
<tr>
<td>rs11883500</td>
<td>0.046</td>
<td>-0.148</td>
<td>0.045</td>
<td>MLPH</td>
<td>T289I</td>
</tr>
<tr>
<td>rs1800422</td>
<td>0.050</td>
<td>0.144</td>
<td>0.050</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs3751107</td>
<td>0.064</td>
<td>-0.137</td>
<td>0.064</td>
<td>MLPH</td>
<td>G172D</td>
</tr>
<tr>
<td>rs3751109</td>
<td>0.084</td>
<td>-0.126</td>
<td>0.084</td>
<td>MLPH</td>
<td>L153P</td>
</tr>
<tr>
<td>rs1052030</td>
<td>0.089</td>
<td>0.125</td>
<td>0.089</td>
<td>MYO7A</td>
<td>T16S</td>
</tr>
</tbody>
</table>
Table 7.19 SNPs that have a significant association with L* in a recessive model (DD vs Dd, dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1800422</td>
<td>0.004</td>
<td>0.212</td>
<td>0.004</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs1058219</td>
<td>0.089</td>
<td>0.125</td>
<td>0.089</td>
<td>MYO5A</td>
<td>R1246C</td>
</tr>
</tbody>
</table>

Tables 7.17 and 7.18 show that rs1800422 is significantly associated with L* in both a dominant and recessive model. The model with the stronger trend (recessive in this case) is more likely to be the correct one. A more obvious example of this in MC1R is shown below (Figure 7.2 and Figure 7.3).

Next, associations with the quantitative variable a* were examined. Table 7.20 shows SNPs where the presence of the minor allele is significantly correlated with a*. Positive correlations indicate an association with hair becoming redder, while negative associations indicate an allelic association with hair that is less red.

Table 7.20 SNPs that have significant allelic associations with a* (D vs. d)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1805007</td>
<td>0.000</td>
<td>0.265</td>
<td>0.000</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs1800422</td>
<td>0.002</td>
<td>0.158</td>
<td>0.002</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs16891982</td>
<td>0.033</td>
<td>-0.110</td>
<td>0.033</td>
<td>DTNBP1</td>
<td>H297Y</td>
</tr>
<tr>
<td>rs11883500</td>
<td>0.041</td>
<td>-0.107</td>
<td>0.041</td>
<td>MLPH</td>
<td>T289I</td>
</tr>
<tr>
<td>rs17470454</td>
<td>0.065</td>
<td>-0.095</td>
<td>0.065</td>
<td>DTNBP1</td>
<td>P272S</td>
</tr>
<tr>
<td>rs3751107</td>
<td>0.076</td>
<td>-0.092</td>
<td>0.076</td>
<td>MLPH</td>
<td>G172D</td>
</tr>
<tr>
<td>rs3751109</td>
<td>0.090</td>
<td>-0.088</td>
<td>0.090</td>
<td>MLPH</td>
<td>L153P</td>
</tr>
</tbody>
</table>

A test for the difference between the three genotypes was conducted. The SNPs shown in Table 7.21 have significant differences of a* between the three possible genotypes of DD, Dd and dd.
Table 7.21 A test for genotypic associations with a* (DD vs. Dd vs. dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1805007</td>
<td>1.96E-10</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs1800422</td>
<td>0.005</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs3751107</td>
<td>0.056</td>
<td>MLPH</td>
<td>G172D</td>
</tr>
<tr>
<td>rs11883500</td>
<td>0.056</td>
<td>MLPH</td>
<td>T289I</td>
</tr>
<tr>
<td>rs3751109</td>
<td>0.072</td>
<td>MLPH</td>
<td>L153P</td>
</tr>
</tbody>
</table>

Table 7.22 and Table 7.23 show significant associations between various SNPs and a* in a dominant and recessive model, respectively. As mentioned above, positive correlations (R) indicate that the minor allele is associated with higher (more red) a* values while negative correlations (R) indicate that the minor allele is associated with lower (less red) a* values.

Table 7.22 SNPs that have a significant association with a* in dominant model (DD, Dd vs dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1805007</td>
<td>0.000</td>
<td>0.277</td>
<td>0.000</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs1800422</td>
<td>0.020</td>
<td>0.171</td>
<td>0.019</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs11883500</td>
<td>0.021</td>
<td>-0.170</td>
<td>0.021</td>
<td>MLPH</td>
<td>T289I</td>
</tr>
<tr>
<td>rs3751107</td>
<td>0.032</td>
<td>-0.159</td>
<td>0.031</td>
<td>MLPH</td>
<td>G172D</td>
</tr>
<tr>
<td>rs3751109</td>
<td>0.040</td>
<td>-0.150</td>
<td>0.039</td>
<td>MLPH</td>
<td>L153P</td>
</tr>
<tr>
<td>rs16891982</td>
<td>0.077</td>
<td>-0.129</td>
<td>0.077</td>
<td>MATP</td>
<td>L374F</td>
</tr>
<tr>
<td>rs17470454</td>
<td>0.080</td>
<td>-0.128</td>
<td>0.080</td>
<td>DTNBP1</td>
<td>P272S</td>
</tr>
</tbody>
</table>

Table 7.23 SNPs that have a significant association with a* in a recessive model (DD vs Dd, dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1805007</td>
<td>0.000</td>
<td>0.430534937</td>
<td>0.000</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs1800422</td>
<td>0.004</td>
<td>0.209</td>
<td>0.004</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
</tbody>
</table>
Tests with $a^*$ show two SNPs that are significantly associated with $a^*$ in both the dominant and recessive model. This happens when phenotypes are not perfectly associated with genotype; however the stronger correlation/trend is likely to be the correct one (recessive in both of these cases). The SNP from MC1R (rs1805007; R151C) is used as an example. The correlation of R151C with $a^*$ is 0.277 in the dominant model and 0.43 in the recessive model. Figure 7.2 shows the slight trend that occurs if the minor allele (D) is treated as a dominant allele (0.277) while Figure 7.3 shows the stronger trend that occurs if D is treated as a recessive allele.

**Figure 7.2** The dominant genotype model for association with $a^*$ for rs1805007 (MC1R; R151C). D represents the minor allele and d represents the major allele.
Finally, associations with the quantitative variable $b^*$ were examined. Table 7.24 shows SNPs where the presence of the minor allele is significantly correlated with $b^*$. Positive correlations indicate an association with hair becoming more yellow, while negative associations indicate an allelic association with hair that is less yellow.
Table 7.24 SNPs that have significant allelic associations with b* (D vs. d)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1805007</td>
<td>0.000</td>
<td>0.214</td>
<td>0.000</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs16891982</td>
<td>0.002</td>
<td>-0.162</td>
<td>0.002</td>
<td>MATP</td>
<td>L374F</td>
</tr>
<tr>
<td>rs1800422</td>
<td>0.004</td>
<td>0.148</td>
<td>0.004</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs7334118</td>
<td>0.018</td>
<td>-0.122</td>
<td>0.018</td>
<td>ATP7B</td>
<td>H1207R</td>
</tr>
<tr>
<td>rs11883500</td>
<td>0.078</td>
<td>-0.092</td>
<td>0.078</td>
<td>MLPH</td>
<td>T289I</td>
</tr>
</tbody>
</table>

A test for the difference between the three genotypes was conducted for b* as well. The SNPs shown in Table 7.25 have significant differences of b* between the three possible genotypes of DD, Dd and dd.

Table 7.25 A test for genotypic associations with b* (DD vs. Dd vs. dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1805007</td>
<td>0.000</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>rs1800422</td>
<td>0.004</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>rs16891982</td>
<td>0.012</td>
<td>MATP</td>
<td>L374F</td>
</tr>
<tr>
<td>rs7334118</td>
<td>0.017</td>
<td>ATP7B</td>
<td>H1207R</td>
</tr>
<tr>
<td>rs3751107</td>
<td>0.044</td>
<td>MLPH</td>
<td>G172D</td>
</tr>
<tr>
<td>rs11883500</td>
<td>0.064</td>
<td>MLPH</td>
<td>T289I</td>
</tr>
<tr>
<td>rs3751109</td>
<td>0.069</td>
<td>MLPH</td>
<td>L153P</td>
</tr>
<tr>
<td>rs729421</td>
<td>0.083</td>
<td>RABGGTA</td>
<td>A420T</td>
</tr>
<tr>
<td>rs1052030</td>
<td>0.086</td>
<td>MYO7A</td>
<td>T16S</td>
</tr>
</tbody>
</table>

Table 7.26 and Table 7.27 show significant associations between various SNPs and b* in a dominant and recessive model, respectively. As mentioned above, positive correlations (R) indicate that the minor allele is associated with higher (more yellow) b* values while negative correlations (R) indicate that the minor allele is associated with lower (less yellow) b* values.
### Table 7.26 SNPs that have a significant association with b* in dominant model (DD, Dd vs dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>RS1805007</td>
<td>0.000</td>
<td>0.261</td>
<td>0.000</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>RS16891982</td>
<td>0.003</td>
<td>-0.215</td>
<td>0.003</td>
<td>MATP</td>
<td>L374F</td>
</tr>
<tr>
<td>RS7334118</td>
<td>0.017</td>
<td>-0.174</td>
<td>0.017</td>
<td>ATP7B</td>
<td>H1207R</td>
</tr>
<tr>
<td>RS11883500</td>
<td>0.034</td>
<td>-0.157</td>
<td>0.033</td>
<td>MLPH</td>
<td>T289I</td>
</tr>
<tr>
<td>RS1800422</td>
<td>0.053</td>
<td>0.142</td>
<td>0.053</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>RS3751107</td>
<td>0.055</td>
<td>-0.142</td>
<td>0.055</td>
<td>MLPH</td>
<td>G172D</td>
</tr>
<tr>
<td>RS3751109</td>
<td>0.081</td>
<td>-0.128</td>
<td>0.081</td>
<td>MLPH</td>
<td>L153P</td>
</tr>
<tr>
<td>RS729421</td>
<td>0.086</td>
<td>-0.126</td>
<td>0.086</td>
<td>RABGGTA</td>
<td>A420T</td>
</tr>
<tr>
<td>RS1052030</td>
<td>0.088</td>
<td>0.125</td>
<td>0.089</td>
<td>MYO7A</td>
<td>T16S</td>
</tr>
</tbody>
</table>

### Table 7.27 SNPs that have a significant association with b* in a recessive model (DD vs Dd, dd)

<table>
<thead>
<tr>
<th>SNP</th>
<th>Corr/Trend P</th>
<th>Corr/Trend R</th>
<th>F-Test P</th>
<th>Gene</th>
<th>Amino Acid Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>RS1805007</td>
<td>0.002</td>
<td>0.230</td>
<td>0.002</td>
<td>MC1R</td>
<td>R151C</td>
</tr>
<tr>
<td>RS1800422</td>
<td>0.002</td>
<td>0.227</td>
<td>0.002</td>
<td>TYR</td>
<td>R402Q</td>
</tr>
<tr>
<td>RS1058219</td>
<td>0.084</td>
<td>0.127</td>
<td>0.084</td>
<td>MYO5A</td>
<td>R1246C</td>
</tr>
</tbody>
</table>
7.3.5 Genetic Interactions

Genetic interactions contributing to Fair/Blonde hair were tested for by performing a regression analysis and looking for second order terms using LL5 Cluster 2 (Fair/Blonde) as the dependant variable (1 = Cluster 2, 0 = Not Cluster 2). The resulting dihybrid genotypes were analysed with the HelixTree Software. Only one significant interaction was found within this sample; between rs16891982 in the MATP gene (L374F) and rs1805005 in MC1R (V60L). The figures below illustrate the significant associations found when the genotypes of rs16891982 and rs1805005 are considered together. Figure 7.4 shows the association of the dihybrid genotypes with Cluster 2 (Fair/Blonde hair) and it can be seen that individuals with genotype GC_GT are all in Cluster 2 (which has a value of 1).

![Figure 7.4](image)

**Figure 7.4** The associations between the rs16891982 and rs1805005 dihybrid and the LL5 Blonde/Fair Cluster 2.

Figure 7.5 and Figure 7.6 show the relationship between the dihybrid genotypes and the quantitative variables L* and b* (a* showed no significant relationship). It should be noted that the genotypes along the x-axis are not always in the same order.
Figure 7.5 The associations between the rs16891982 and rs1805005 dihybrid and L* value

Figure 7.6 The associations between the rs16891982 and rs1805005 dihybrid and b* value
7.4 Discussion

7.4.1 Associations with Categorical Variables (Clusters)

In this analysis, five SNPs show various significant associations, in this population, with the hair colours as defined by the cluster analysis discussed in Chapter 3. These associations will be discussed; noting that D represents the minor allele and d represents the major allele found in this population. Since all SNPs in this study were non-synonymous mutations, the SNP will be referred to by the amino acid substitution listed in the Entrez SNP database [97].

As expected, a SNP from the MC1R gene (R151C) is associated with Red hair (Cluster 1) at the allelic level, as is a SNP from the TYR gene (R402Q). The high odds ratio of this SNP in MC1R (13.1) indicates the strength of this relationship. MC1R R151C is well known as a highly penetrant red-hair allele [14].

There are allelic associations with Fair/Blond hair in the genes DTNBP1 (H297Y) and MYO7A (C1628S). There are also dominant and recessive genotype associations with these two SNPs, as well as in MC1R (R151C), TYR (R402Q) and ATP7B (R832K).

When Cluster 1 (Red) is removed from the sample, the SNPs previously associated with Red hair are now associated with Fair/Blonde hair; however the odds ratios are much weaker (i.e. 2.3 compared to 13.1 for rs1805007). R832K in ATP7B is no longer associated with Cluster 2 when Cluster 1 is removed. The strongest associations (highest odds ratios) found within this sample were for H297Y in DNTBP1 in a dominant model (DD/Dd vs. dd) with an OR of 8.0 (when Reds were included in the sample) and R402Q in TYR in a recessive model with an OR of 3.8 (when Reds were excluded from the sample).

Associations were also found with Cluster 5 (Dark hair) in this sample. A SNP in the gene MATP (L374F) was significantly associated with dark hair in both the allelic and dominant model, which corresponds to previously published work on this gene [45]. Two SNPs in ATP7B (S406A and V456L) were also found to be associated in a
dominant model with Cluster 5 (Dark hair). The odds ratio for these two SNPs was less than one (0.49); therefore it is the major allele (d) that is associated with dark hair for these two polymorphisms and the 406S and 456V amino acid status. It should be noted that these are different SNPs than were associated with Blonde/Fair hair (that was R832K) and not just opposite alleles.

Of the six genes found to be significantly associated with hair colour clusters, several are familiar genes in the pigmentation pathway and are mentioned in Section 7.1.1. TYR codes for the rate-limiting melanogenesis protein, Tyrosinase [86], while MC1R (the melanocortin 1 receptor) is primarily involved with the switch between eumelanin and pheomelanin [4, 14, 86, 90]. MATP (membrane associated transport protein) is also referred to as SLC45A2, a solute carrier and a melanocyte differentiation antigen, thought to be involved in the mediation of melanin synthesis [45, 97].

Less familiar genes include DTNBP1, the gene for dystrobrevin-binding protein 1, which has been found to be associated with human skin pigmentation [109] and is thought to be involved in organelle biogenesis, including that of melanosomes [97]. The H297Y polymorphism in DTNBP1 was quite rare in this population, with a minor allele (A) frequency of 0.008 and this may contribute to the strong effect seen for this allele.

The gene ATP7B also appears to be associated with hair colour clusters. This gene codes for a Copper transporting ATPase [97, 110] and has been implicated in pigmentation of *Drosophila melanogaster* (fruit flies) [110]. The ATP7B protein is thought to control the exit of copper from the cells (Wilson disease in humans is caused by a defective ATP7B protein and results in toxic copper accumulation) [110]. Tyrosinase is a copper-dependant enzyme [111] and therefore; variations in cellular copper caused by polymorphisms in ATP7B may affect Tyrosinase activity.

The gene MYO7A codes for a Myosin VIIA protein, a mechanochemical protein that, in complex with Rab27a, is involved with the movement and distribution of melanosomes in the retinal pigmented epithelium [112] and may have similar effects in other pigmented tissues, such as hair.
7.4.2 Associations with Quantitative Variables (L*, a* and b*)

Similar results to Section 7.4.1 were obtained when tests for association with the quantitative variables L*, a* and b* were performed with various models – allelic, genotypic, dominant and recessive.

Polymorphisms in the genes TYR, MATP, DTNB1, ATRN, MC1R, MLPH, MYO7A and MYO5A were found to be associated with L* values and these results tend to correspond with genes that were associated with light or dark hair in Section 7.4.1. For example, polymorphisms in MATP that were associated with Dark Cluster 5 have a negative correlation to L*, as would be expected. Seven new polymorphisms (not associated in the categorical analysis) were significantly associated with L* in at least one of the models and these include two SNPs in the gene ATRN (I426T and R1152K), three SNPs in the gene MLPH (G172D, L153P and T289I), a different SNP in MYO7A (T16S) and one in MYO5A (R1246C).

The ATRN gene codes for the attractin protein; the mouse ortholog of this protein (mahogany) is involved in the melanocortin 1 receptor pathway and therefore may have an effect on the pheomelanin production [113].

The melanophilin protein is coded by MLPH and forms a motor complex with Rab27a and Myosin 5A (MYO5A) that has been found in mice to tether the melanosomes to the cytoskeleton for movement around the cell [94].

Tests for associations with a* achieved very similar results to those with L* with two notable exceptions. R151C in MC1R shows a much higher correlation (R=0.43) with a* compared to other correlations in this sample set; indicating that it is a strong predictor of redness in hair, as expected. A second SNP in DTNB1 (P272S) shows a small negative correlation with a*.

The same polymorphisms continue to be associated with b*, with the inclusion of H1207R in ATP7B and A420T in RABGGTA.
RABGGTA is a Rab geranylgeranyl transferase \( \alpha \) sub-unit that is involved in the prenylation (adding lipids) of a Rab protein and is thought to anchor it to a membrane [114] and therefore is somewhat involved in the transfer of melanosomes.

### 7.4.3 Genetic Interactions

Two polymorphisms, one in the MATP gene (L374F) and one in MC1R (V60L) were shown to interact in their contribution to Fair/Blonde hair (they are associated with LL5 Cluster 2). It is possible that MATP;L374F may act as a suppressor of the fair hair in MC1R;V60L individuals, however many more samples would need to be analysed to elucidate the cause of any possible interactions. V60L is a different SNP in MC1R than is associated with red hair colour and is known as a low-penetrance red hair allele [14, 96].

### 7.4.4 Conclusions

It is clear that even in this small sample, there are significant associations with hair colour as defined by the cluster analysis in Chapter 3, particularly associations with Red, Fair/Blonde and Dark hair. There are also many significant correlations between genotypes at various SNPs and the quantitative measurements of \( L^* \), \( a^* \) and \( b^* \). As expected, one polymorphism or even one gene does not provide all of the information about hair colour and many genes and multiple polymorphisms within various genes seem to have an effect on hair colour. Most of these genes are familiar to those studying pigmentation; however, several are novel in the context of human hair pigmentation, though all have a logical connection to the pigmentation pathway. An expansion of the sample size will aid in further examination of these polymorphisms and may clarify and strengthen the associations found. An increase in the number of samples that were examined microscopically and chemically would allow for the further testing of genetic associations with these features, independently of their macroscopic colour values and clusters. With the current number of samples that have been microscopically and chemically analysed, no associations or trends were observed (data not shown). Additional SNPs should also be considered in the future. Several other
SNPs have recently been found by others to be associated with self-reported ‘Blonde’ hair [46], including additional SNPs in the genes SLC24A4, KITLG, TYR and OCA2 that have not been tested in this population or in another objectively measured one.

Another important next step in this work would be to conduct blind population assignment tests, to determine if likely genotype combinations can correctly assign individuals to a particular group or cluster. This approach would further validate results of the genetic associations and advance the practical applications of this type of work.
Chapter 8  Discussion and Conclusions

8.1  Summary

It was hypothesised in Chapter 1 that there are genetic variants specifically associated with an objectively defined Blonde phenotype. This project was conducted with an aim to first characterise and define the phenotype under study (‘blonde hair’) and then to evaluate DNA sequence variants that may be associated with this phenotype. These variants may be useful in predicting the hair colour of an unknown subject in forensic cases where DNA is found at a crime scene, but there is no suspect sample to compare it to.

The previous chapters have examined hair colour at several levels in accordance with the project aims. Macroscopic colour was measured with reflective spectrophotometry and digital image analysis, followed by a microscopic analysis of hair colour and pigmentation pattern. Next, a chemical characterisation was carried out, followed by analysis of genetic variations within genes thought to be involved in the pigmentation process.

Chapter 3 investigated the measurement of macroscopic colour by reflective spectrophotometry. This analysis showed that while hair colour can be perceived as a continuum, it can be divided into categories or clusters based on spectrophotometric measurements in the CIE L*a*b* colour space and these clusters can be well discriminated from each other. The importance of the objective measurement of individuals to be included in research studies is indicated by the discrepancies between self-reported hair colour and observer-reported hair colour and the poor separation of observer-reported colour groups compared to those defined by clustering analysis. The most favourable cluster grouping achieved in this analysis, for the purpose of this study, is called LL5 - when the log likelihood method is used to break individuals into five hair colour groupings, named Dark, Medium, Fair, Red and White.

Chapter 4 examined the use of digital image analysis to measure hair colour. This analysis showed that although measuring colour from digital images is convenient, the
colours measured and classified do not correspond very well to those colours measured by reflective spectrophotometry and would have limited use in studies requiring accurate measurement of hair colour, such as genetic association studies. They also do not provide an accurate description of hair colour for crime investigators to utilise as visual identifiers or in description announcements.

The microscopic colour of hair was examined in Chapter 5 and from this analysis, it can be seen that there is a strong relationship between macroscopic and microscopic $L^*$ values that would be useful if one was trying to predict the colour of a head of hair (macroscopic) from a strand (microscopic). The relationship between hair lightness ($L^*$) and thickness is also clearly illustrated at both the macroscopic and microscopic levels.

The results of the chemical characterisation described in Chapter 6 show how the amount of eumelanin in hair (represented by its degradation product – PTCA) is related to the colour as measured in the CIE $L^*a^*b^*$ colour space by reflective spectrophotometry and microscopic analysis. As would be expected, lighter hair has less eumelanin; however, there is a stronger relationship between the amount of eumelanin present and the value of the $b^*$ (yellow) axis than with the $L^*$ or $a^*$ axes. The amount of PTCA measured is more closely related to the microscopic colour than to the macroscopic colour, which may be due to the different methods of colour measurement (reflective spectrophotometry compared to a digital microscopic image). The PTCA results obtained for the LL5 clusters are also as expected. Clusters 5 (dark), 4 (Medium) and 2 (Fair) have progressively less of the eumelanin degradation product. The quantification of the pheomelanin degradation product (AHP) was unsuccessful in this sample in the time available for analysis.

Chapter 7 involves the genetic analysis with regards to both categorical variables - the LL5 clusters - and the quantitative variables – the $L^*$, $a^*$ and $b^*$ colour values. In the categorical analysis, five of 58 SNPs show various significant associations with Fair/Blonde hair (Cluster 2). These SNPs are in the genes MC1R (R151C), DTNBP1 (H297Y), TYR (R402Q), ATP7B (R832K) and MYO7A (C1628S). MC1R (R151C) and TYR (R402Q) were found to be associated with Red hair (Cluster 1) and associations were also found with Cluster 5 (Dark hair) in this sample. A SNP in the
gene MATP (L374F) was significantly associated with dark hair in both the allelic and dominant model, which corresponds to previously published work on this gene [45]. Two SNPs in ATP7B (S406A and V456L) were also found to be associated in a dominant model with Cluster 5 (Dark hair).

In the quantitative variable analysis, polymorphisms in the genes TYR, MATP, DTNBP1, ATRN, MC1R, MLPH, MYO7A and MYO5A were found to be associated with L* values and these results tend to correspond with genes that were associated with light or dark hair in the categorical analysis. Seven new polymorphisms (not associated in the categorical analysis) were significantly associated with L* in at least one of the models and these include two SNPS in the gene ATRN (I426T and R1152K), three SNPs in the gene MLPH (G172D, L153P and T289I), a different SNP in MYO7A (T16S) and one in MYO5A (R1246C).

Tests for associations with a* achieved very similar results to those with L* with two notable exceptions. R151C in MC1R shows a very high correlation (R=0.43) with a* compared to other correlations in this sample set; indicating that it is a strong predictor of redness in hair, as expected. A second SNP in DTNBP1 (P272S) shows a small negative correlation with a*. The same polymorphisms continue to be associated with b*, with the inclusion of H1207R in ATP7B and A420T in RABGGTA.

Two polymorphisms, one in the MATP gene (L374F) and one in MC1R (V60L) were shown to interact in their contribution to Fair/Blonde hair (they are associated with LL5 Cluster 2).

### 8.2 Discussion

It was hypothesised in Chapter 1 that there were genetic variants specifically associated with and objectively defined Blonde phenotype. To test this hypothesis, one must first define the phenotype under question and then search for any genetic associations. With this in mind, there were three primary objectives for this research project (listed in Chapter 1). The first objective was to examine the natural, adult hair colour variation among a population of European ancestry and to evaluate methods for both the
measurement and classification of hair colours – what is the phenotype and what are its limits?

The results of Chapters 3 and 4 fulfilled this objective very well. Natural, adult hair colour was studied extensively and the fair-to-dark hair colour spectrum has been characterised more thoroughly than previously reported in the literature [52, 59]. Blonde or Fair hair colour can be defined in several ways, by self or observer-reported colours or by various statistical clustering analyses using measurements such as reflective spectrophotometry or digital image analysis. This study found that objective measurements taken by reflective spectrophotometry and subject to statistical cluster analysis provided groupings that were objective, describable and useful as a phenotype due to the resemblance these groupings had to our subjective descriptions of hair colour. Even if the same clusters are not found in future samples (as is expected given changes in sample size and in various populations), the method can be used to achieve objectivity in classifying hair colours and is more likely to lead to an easier description of individuals and comparisons between research studies. If a large standardised database was to be constructed, particular groupings could be defined and then used by researchers to achieve consistency in their investigations. Such a database could also be used by forensic investigators and police to achieve consistent descriptions of hair colour.

There are a few other studies that have objectively measured hair colour [41, 52, 59] and having this data allows for potential comparisons between phenotypes and results of genetic studies. However; previous studies did not examine hair colour to the extent done here. As reported in Chapter 3 (Section 3.1.2), hair colour has been measured by reflective spectrophotometry; however these studies have used only the L* axis or the M index [59] or have concentrated on red hair [41, 52]. One study carefully measured hair colour (including red and blonde hair in particular) by reflective spectrophotometry, but used subjects of 12 and 14 years of age [41], running the risk of the subjects hair colour changing due to hormonal influences during puberty, such as the darkening of blonde hair [51].

Being able to measure hair colour from digital photographs may assist in efforts to construct or expand such a database and may also assist in various other areas with an
interest in hair colour. This may speculatively include fields such as surveillance, cosmetics or others. In this study, hair colour measured by digital image analysis did not correspond well to the reflective spectrophotometry standard and showed a high amount of variation, however; it may be possible for improvements in technology or software to change this outcome.

There are, of course, problems in the measurement of hair colour and in the future use of the data. While all volunteers in this study were healthy young adults with their natural hair colour, it is always possible that hair colours may be changed due to non-genetic influences such as sunlight, hormonal changes and other epigenetic factors and the use of hair products [16, 17, 51]. While every effort was made to reduce the influence of these processes for this early-stage research, they will have effects on hair colour in practice and further research into each of these factors may assist in the characterisation of hair colours, which will eventually assist in any use of predictive forensic DNA tests or the use of hair colour data in traditional hair examinations. In addition, not all DNA samples tested will belong to young healthy adults and so further research into hair colour changes due to age or medical conditions may also be worthwhile pursuing.

The second objective was to characterise the microscopic and chemical features of hair colour in this sample and how they relate to the macroscopic colour – what are the properties of this phenotype? The microscopic analysis undertaken for this project (Chapter 5) provided some very interesting, new results. While microscopic hair colour has been examined to a small extent, in an effort to assist with traditional hair examinations and identification of hairs [24, 27], there seem to be no previous studies relating the microscopic hair colour to the macroscopic, a deficit in the literature that this research will help fill. Traditional forensic hair examiners often offer an opinion to the hair colour of an individual based on a microscopic analysis of a strand of hair [22, 115]. Having statistical data to support such claims will assist in this type of examination. It may also help link the various levels of analysis – genetic, chemical, microscopic and macroscopic – providing information to the scientist trying to predict hair colour based on any of the above data (particularly genetic). Due to the significant results achieved in this experiment, it would be beneficial to continue the analysis on an expanded sample size. The examination of more individuals and more hairs from each
individual would assist in the further analysis of the relationship between microscopic and macroscopic hair colour.

The chemical analysis in Chapter 6 supported previous research by finding similar eumelanin levels in various hair colour categories, as discussed in Section 6.4 [20, 82], but having the macroscopic and microscopic hair colour measurements allowed for a more specific relationship to be established between the amount of PTCA recovered (the eumelanin degradation product) and the colour of the hair. Chemically characterising more of the individuals in this sample or on a different, expanded sample will further elucidate relationships between pigment composition and colour. It is widely accepted that less eumelanin pigment leads to a lighter hair colour and this experiment does not show anything unexpected. It does, however, provide data for this particular sample and it provides chemical characterisation of the reflective spectrophotometry clusters. This may be one way for various researchers to compare their work, if colour measurements were not taken. If enough samples could be collected for chemical analysis, it would be worthwhile conducting genetic association tests directly on the amount of pigment present in the hair samples. Unfortunately, the 55 individuals in this sample with chemical data do not provide enough power for meaningful genetic analysis but should be reconsidered in the future.

It would also be beneficial to obtain pheomelanin characterisation data for this sample. Even though eumelanin is the primary pigment of interest when studying the light-to-dark continuum of hair colour, the ratio between the eumelanin and pheomelanin pigments is relevant to the study of hair colour genetics. In the time available for this analysis in the current study, the analysis of the pheomelanin pigments could not be made to work. Further efforts would eventually change this outcome and the data could be incorporated into the analysis.

The third and final objective was to compare genetic results with an objectively measured and defined hair colour phenotype in the search for genetic associations with ‘blonde’ hair colour. This was a small, but well-characterised sample and as reported in Chapter 7, several significant genetic associations were found with Fair/Blonde hair as defined in this cluster analysis. These results support other genetic studies that have been released recently [45, 46] by finding associations in some of the same genes (such
as MC1R and TYR), with the small effects expected in the genetic analysis of hair colour [9]. This study also offered some new associations that would be worth examining in the future on a larger sample. The genes that showed significant associations in this study were in various areas of the pigmentation pathway, including the manufacture of melanin (melanogenesis) as expected, but many were involved in other areas of pigmentation, such as the formation of melanosomes and the movement of melanosomes around the cell. These genes seem to be more involved in the light-to-dark differences in pigmentation within individuals of European ancestry than those genes typically involved in population differences (melanogenesis [4, 34]) and red hair (pheomelanin switch [4, 34]). Looking at other genes involved in melanosome biogenesis and movement and examining the effect of causal variations in these genes would be an important part of future research into genetics and cellular biology of pigmentation. As well as examining additional candidate genes, looking for associations with clusters or quantitative colour measurements using whole genome scans may provide additional information in the search for hair colour genes.

There is a great deal of work left before the prediction of hair colour from genetic information is remotely possible, however this study will provide future researchers with associations that can be expected in an objectively measured population. Large, objectively measured samples are difficult to come by. If developments in technology allow for the easier measurements of hair colour (by any method), this will become more likely and genetic analysis may become more accurate. Without objective measurements, the prediction of physical features for forensic prediction is still possible; however it will be less accurate and may be open to more challenges from investigators and the justice system.

The results of this study address some of the issues in human hair colour genetics that were raised in Section 2.3.3. This study provides additional evidence that the genes involved with hair colour are somewhat different to those most strongly associated with eye colour and that there is a larger number of gene variants with lower penetrance involved in hair colour. 19 SNPs were found to have some association with hair colour in this study and most of these SNPs had fairly small effects (odds ratios of 1 to 4 or correlations between 0.1 and 0.3), a trend which was also seen in the deCODE study mentioned previously [46]. As well, OCA2 (previously shown to be the main eye
colour gene [116]) does not seem to have a strong enough effect on hair colour to be seen in this sample, if any, supporting the idea that different genes are involved.

There is also some indication that epistatic genetic interaction may contribute to differences in hair colour, such as the interaction found between MATP and MC1R in this study and a previous interaction between MC1R and a SNP in the 3’ untranslated region of the ASIP gene [44]. Although one interaction was detected in this sample, due to generally low gene effects, the study population would likely have to be much larger to find any more such interactions.

One of the issues in finding genetic associations with hair colour that has been mentioned in the literature [9] is that the phenotyping of hair may not be ‘specific’ enough to find consistent and specific associations. While this still may be true, to some degree, this project has provided a great deal of information on objectively defining hair colour phenotypes and offers methods and data for the future description of hair colours in various types of research studies.

8.3 Future Considerations

As a pilot study, the presented research has raised some interesting ideas that would benefit from further examination. The most important consideration for future research would be to expand the sample size. All results obtained in this study are statistically significant; however, an increase in the sample size can only improve the power of analysis, particularly for the genetic analysis. An increase in the number of individuals examined by microscopic and chemical analysis would also benefit further studies by more strongly illustrating existing relationships. Increasing the number of individuals with objectively measured hair colour would assist in the development of a ‘phenotypic database’ – a database with images of all possible hair colours. Sample sizes consisting of thousands of individuals (such as in the deCODE genetics study discussed in Chapters 2 and 7) will eventually be necessary for both hair colour research and genetic association tests and the challenge of objectively measuring the hair colour of that many individuals will take a lot of work. The use of more modern reflective spectrophotometers (that can output results directly to a notebook computer) would
assist in this effort and do exist, but were not available for this study. The development of digital photographic equipment and colour measurement software specific to measuring hair colour may also be helpful.

Another area of research that would be useful and interesting is the further study of the cellular biology of hair pigmentation. One study by Commo et al. (2004) examined the distributions of different proteins in African, Asian and Caucasian hair bulbs and as a part of their study, tagged the pMel-17 melanosome identifier protein [50]. Since some of the genetic variations associated with blonde hair are in genes that control the biogenesis and movement of melanosomes, it may be worth using a similar methodology to examine the distribution of these organelles in hair bulbs within a population of European ancestry across various hair colours to determine if the number or distribution of melanosomes is correlated with hair colour. This would increase the understanding of the underlying pigmentation processes and would further elucidate the specific genetic contributions to hair colour.

Further examination of the exact composition of hair melanin pigments would be another area of research that would assist in the characterisation of hair for forensic or genetic purposes. While perhaps unlikely, improved availability of and information on melanin degradation product standards (PTCA, PDCA and AHP) would assist in method development and allow for more thorough analysis.

Another logical progression from this study would be the genotyping of several SNPs that have been associated with fair or blonde hair colour in the literature published since this set of 58 SNPs was chosen. These include SNPs in the genes SLC24A4, KITLG, TYR, OCA2 and MC1R [46] and possibly in other genes implicated. It may also be worth looking at more SNPs in genes found to have significant associations in this study, there may be a haplotype effect to be considered or these SNPs may be in linkage disequilibrium with the causative mutation. As well as finding significant associations, discovering how these causative mutations affect proteins produced and how they may interact with other genes will help researchers understand hair colour and will lead to more accurate predictions.
8.4 **Conclusion**

In conclusion, this study successfully met its aims of objectively classifying and characterising blonde (or fair) hair colour and finding genetic associations with blonde hair colour. Some of the data supports previous studies in the area of hair colour characterisation and genetics and a great deal of it is new and can be useful to researchers in forensic science and in other areas of pigmentation research. There are several matters that may be expanded on in the future and it is hoped that this study may be a resource to those conducting hair pigmentation research in anthropology, genetics and in forensic science.

*Colours are the smiles of nature.*

--*Leigh Hunt (1784-1859)*
Chapter 9

References


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APPENDIX 1 – Raw Data
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</tbody>
</table>

1. Observer Reported Colours: 1 = Blonde  2 = Light Brown  3 = Dark Brown  4 = Black  5 = Red  6 = White
2. Non-European Individuals
APPENDIX 2 – Computer Code
aRGB2LAB Conversion Module for V++

{Images need to be .jpg in the following folders:
- folder containing subfolders:
  - images
  - white balance images
The images for testing are saved to the 'images' folder.
The white balance images are saved in the 'white balance images' folder.
The name of the 'white balance image' must be the same as the 'image' with a 'w' appended as a suffix.}

button btn_text,'Lab stats from folder w/ white correction';
var
img,ref_present, white_ref;
temp_small,temp_big;
img_red,img_green,img_blue;
ref_red,ref_green,ref_blue;
l,la,lb;
rl,ra,rb;
t_small, t_big;
name, imgname, whiterefname, path;
ed,ed2;

const                        //**************************************************
white level values
white_l=93.4;
white_a=1.16;
white_b=-2.06;

function BrowseFolders( Path:pointer ) : integer ; external
'VppCommonDlgs.dll';

function GetFolder ;
var
FolderName ;
begin
FolderName := StringOfChar( ' ',255  ) ;
if BrowseFolders( FolderName ) <> 0 then
  GetFolder := Trim( FolderName )
else
  GetFolder := '' ;
end;   GetFolder }

function powerof(s,t);//*********************************** get 's' to
the power 't'
var
value,result;
begin
value:=Ln(s);
result:=Exp(value*t);
powerof:=result;
end;

function get_channel(i);//********************************* get RGB
channel
var
t_small,t_big,c;
begin
c:=single(i)/255;
t_small:=c*(c<=0.04045)/12.92;
t_big:=powerof(((c+0.055)/(1.055),2.4)*(c>0.04045));
get_channel:=t_small+t_big;
function get_l (i,r,g,b);//************************************** get L channel
var
yn,y,yy,temp_big,temp_small;
begin
yn:=0.212671+0.715160+0.072169;
y:=0.212671*r+0.715160*g+0.072169*b;
yy:=y/yn;
temp_big:=( 116*powerof(yy,0.3333) -16)*(yy>0.008856);
temp_small:=yy*(yy<=0.008856)*903;
get_l:=temp_big+temp_small;
end;

function get_a (i,r,g,b);//************************************** get a channel
var
xn,yn,x,xx,yy,temp_big,temp_small;
begin
xn:=0.412453+0.357580+0.180423;
y:=0.212671*r+0.715160*g+0.072169*b;
x:=0.412453*x+0.357580*g+0.18042*b;
xx:=x/xn;
y:=y/yn;
temp_big:=powerof(xx,0.3333)*(xx>0.008856);
temp_small:=(7.787*xx+0.1379310345)*(xx<=0.008858);
xx:=temp_big+temp_small;
temp_big:=powerof(yy,0.3333)*(yy>0.008856);
temp_small:=(7.787*yy + 0.1379310345)*(yy<=0.008856);
yy:=temp_big+temp_small;
get_a := 500*(xx-yy);
end;

function get_b (i,r,g,b);//************************************** get b channel
var
yn,zn,y,z,yy,zz,temp_big,temp_small;
begin
yn:=0.212671+0.715160+0.072169;
zn:=0.019334+0.119193+0.950227;
y:=0.212671*r+0.715160*g+0.072169*b;
z:=0.019334*r+0.119193*g+0.950227*b;
yy:=y/yn;
zz:=z/zn;
temp_big:=powerof(yy,0.3333)*(yy>0.008856);
temp_small:=(7.787*yy + 0.1379310345)*(yy<=0.008856);
yy:=temp_big+temp_small;
temp_big := powerof(zz,0.3333)*(zz>0.008856);
temp_small := (7.787*zz + 0.1379310345)*(zz<=0.008856);
zz := temp_big + temp_small;
get_b := 200*(yy-zz);
end;

begin//******************************************************* program start
path:=getfolder;
Name := FindFirstFile( path+'\images\*.jpg',fa_Archive ) ;
ed:=createeditor('output');
ed2:=createeditor('corrected mean values');
writeln(ed2,chr(9)+'L'+chr(9)+'a'+chr(9)+'b');
while StrLen( Name ) > 0 do
begin
  imgname:=name;
  open(imgname,img);
  converttype(img,typ_rgbfloat);
  img:=img+0.01;
  //writeln(imgname);
  whiterefname := path+'\white balance images\'+w+extractfilename(imgname);
  open(whiterefname,white_ref);
  //writeln(whiterefname);
  img_red:=get_channel(red(img));
  img_green:=get_channel(green(img));
  img_blue:=get_channel(blue(img));

  l:=get_l(img,img_red,img_green,img_blue);//***************************
  ****************
  get image Lab channels
  la:=get_a(img,img_red,img_green,img_blue);
  lb:=get_b(img,img_red,img_green,img_blue); //***************************
  //********************************************************* write image details
  writeln(ed,'____________________________________');
  writeln(ed,'Image for conversion is:' );
  writeln(ed,extractfilename(imgname)+chr(9)+'L'+chr(9)+'a'+chr(9)+'b');
  writeln(ed,'Mean'+chr(9)+str(meanof(l))+chr(9)+str(meanof(la))+chr(9)+str(meanof(lb)));
  writeln(ed,'Deviation'+chr(9)+str(stdof(l))+chr(9)+str(stdof(la))+chr(9)+str(stdof(lb)));
  writeln(ed);
  //********************************************************* get white reference Lab channels
  ref_red:=get_channel(red(white_ref));
  ref_green:=get_channel(green(white_ref));
  ref_blue:=get_channel(blue(white_ref));

  rl:=get_l(white_ref,ref_red,ref_green,ref_blue); //show(rl);
  ra:=get_a(white_ref,ref_red,ref_green,ref_blue); //show(ra);
  rb:=get_b(white_ref,ref_red,ref_green,ref_blue); //show(rb);
  //********************************************************* write whitebalance image details
  writeln(ed,'White reference image is:' );
  writeln(ed,extractfilename(whiterefname)+chr(9)+'L'+chr(9)+'a'+chr(9)+'b');
  writeln(ed,'Mean'+chr(9)+str(meanof(rl))+chr(9)+str(meanof(ra))+chr(9)+str(meanof(rb)));
  writeln(ed,'Deviation'+chr(9)+str(stdof(rl))+chr(9)+str(stdof(ra))+chr(9)+str(stdof(rb)));
  writeln(ed);
  //********************************************************* write corrected image details
  writeln(ed,'Corrected image values:'); //*************************** write corrected image details
  writeln(ed,'Mean'+chr(9)+str((white_l-meanof(rl))+meanof(l))+chr(9)
  +str((white_a-meanof(ra))+meanof(la))+chr(9) +str((white_b-
  meanof(rb))+meanof(lb)));
Name := FindNextFile;
end;

copytoclipboard(ed2);
writeinfo('complete');
free(img);
free(l);
free(la);
free(lb);
free(rl);
free(ra);
free(rb);
end
APPENDIX 3 – Project Information Sheet, Consent Forms and Questionnaire
Blond Hair Colour: Classification and Genetic Associations

Background Information

This study is conducted at Victoria University by PhD candidate, Michelle Vaughn and her supervisor, Dr. Swati Baindur-Hudson (03 9216 8123) with the support of Dr. Roland van Oorschot (Associate Investigator and Co-supervisor) from the Victoria Police Forensic Services.

Researchers all over the world are developing tests for physical features such as eye colour and red hair. This study is concerned with characterising hair colour and looking at genetic variations that may contribute to blonde or fair hair. To do this we need to examine the hair at many levels and look at a persons DNA to find the correlations.

If the research is successful, the ability to describe an individual from their DNA has many potential uses:

It may be possible in the future to determine some of a person’s physical features from DNA inadvertently left at a crime scene. This would allow the police to use this information as an investigative tool for limiting a suspect pool, or to provide a starting point for a search, much as geographical or psychological profiling are used now in many jurisdictions. Other circumstances where it would be useful to be able to predict the physical features of a person from their DNA include natural and criminal mass disasters, where the appearance of victims may have been destroyed beyond recognition. If DNA could be recovered from these victims and some of their physical features predicted it may make it easier for forensic investigators to narrow the search for their identity.

The procedures and samples required for this study are listed on the back of this information sheet. You may choose to participate in some or all of them, at your discretion.

While this research may one day benefit forensic scientists and investigators, your samples are going to be used purely for research, your identity kept anonymous, and information about you or your DNA sample will kept totally confidential.

Steps taken to ensure privacy, anonymity and confidentiality of volunteers

Ethics approval has been obtained from the Victoria University Human Research Ethics Committee, approval number HRETH 06/156. Ethics approval is necessary to protect the rights of the people taking part in this study. The conditions of the approval include:

- Samples will only be taken from volunteers with informed consent.
- Each volunteer has the opportunity to withdraw from the project at anytime.
- All details recorded will be kept in strictest confidence, under lock and key and only the researchers will have access to any information given while respecting the requirements of the law.
- Samples and questionnaire answers will only be identified by code number not by name.
- To ensure the privacy of volunteers, data will only be presented as linked to physical characteristics such as hair colour, and not related to individuals
- Neither your name nor contact details will be linked to samples or data generated from analysis. The data will only be linked to physical characteristics.
- All names and contact details will be destroyed at the end of the study
- At the completion of the entire study, all details as well as any remaining swabs and extracted DNA samples will be destroyed.

To ensure your safety, privacy and confidentiality at all times, we have taken a number of measures:

Physical: The sampling method used will be the collection of inner cheek cells with sterile swabs, which will only be taken by you. This will make the procedure quick, easy and least invasive for you. If hair is to be plucked and clipped, all efforts will be made to reduce discomfort.

Psychological/social/legal: Any data collected from the answers to the questionnaire or from the analysis of samples will be used for research purposes only and will not be made available to any other party. Strict confidentiality and security measures will be maintained – see details above.
Procedures:

- After reading the above information, you will be asked if you want to participate in the study.

- If so, you will be asked to fill out and sign the consent form

- You will be assigned a code and from there on your samples will only be referred to with the code to preserve your anonymity.

- You will then be asked to provide a sample of inner cheek cells (buccal cells) using sterile swabs.

- If you have given permission, you will be requested to fill out a questionnaire about some of your physical characteristics, which will only be identified by your assigned code, no personal details

- If have given permission, a digital photo will be taken of your hair for colour determination. Measurements of your hair colour will also be taken with a reflective spectrophotometer. This instrument uses normal light to measure colour. Again, all these will be recorded against the code only.

- If you have given permission, several hairs will be plucked from your scalp and approximately 30 mg (a small amount) of hair will be clipped.

- The collected samples will be stored and may be used to study links between genetic variations and the physical characteristics of hair colour.
Adult Consent Form

I, __________________________________________________________________________ of
(address) ______________________________________________________________________

 certify that I am at least 18 years old and that I am voluntarily giving my consent to
participate in the experiment entitled:

Blonde Hair Colour: Classification and Genetic Associations

being conducted at Victoria University, by PhD candidate, Michelle Vaughn and her supervisor, Dr.
Swati Baimdur-Hudson, with the support of Dr. Roland van Oorschot (Associate Investigator and Co-
supervisor) at the Victoria Police Forensic Services.

I certify that the objectives of the experiment, together with any risks associated with the procedures to
be carried out in the experiment, have been fully explained to me by

(Name of researcher) __________________________________________ on (date) ____________

(Signature of researcher): ________________________________

and that I have had the opportunity to have any questions answered. I also understand that I can
withdraw from this experiment at any time and that this withdrawal will not jeopardise me in any way. I
have been informed that the information I provide will be kept confidential.

I freely consent to participation in the following procedures:

Having a digital photo and other colour measurements taken of myself to help determine the exact
colour of my hair……………………………………………………………………………… ☐Yes ☐No

Filling out of a questionnaire regarding some of my physical characteristics……………… ☐Yes ☐No

Self-taking of buccal cell samples (for DNA) using sterile swabs……………………… ☐Yes ☐No

Plucking of several scalp hairs…………………………………………………………… ☐Yes ☐No

Trimming of approximately 30 mg of scalp hair ………………………………………… ☐Yes ☐No

To be contacted if need be in future, should a sample need to be repeated……………… ☐Yes ☐No

If yes, your telephone no.: _______________ and/or email address: ______________________

Signature: ________________________________ Date: ____________________
Revocation of Consent Form

I, _________________________________________________ of
(address) ________________________________________________

___________________________________________________________

hereby wish to WITHDRAW my consent to participate in the research project

Blonde Hair Colour: Classification and Genetic Associations

being conducted at Victoria University, by PhD candidate, Michelle Vaughn and her supervisor, Dr. Swati Baindur-Hudson, with the support of Dr. Roland van Oorschot (Associate Investigator and Co-supervisor) at the Victoria Police Forensic Services.

I understand that such withdrawal WILL NOT jeopardise any treatment or my relationship with Victoria University or the Victoria Police Forensic Science Services (if applicable)

Any data already collected may / may not be included in the research project (circle one).

Signature ____________________________________________ Date: ________________
**General Information**

1. **Reason for interest in project:** 
2. **Date of Birth:** _____ / _____ / _____
   (Day / Month / Year)
3. **Gender:** Male □ Female □
4. **Do you currently have your natural adult hair colour?** □ Yes □ No
5. **Are you taking any medication or nutritional supplements that have altered your adult hair colour?** □ Yes □ No

**Pigmentation Characteristics**

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<th>Dark Brown □</th>
<th>Light Brown □</th>
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<td>Blue □</td>
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Or describe in your own words:

**Background** *(please tick appropriate box for each)*

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<th>Maternal Grandfather</th>
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</tbody>
</table>

* North European: France, Switzerland, Belgium, Netherlands, Denmark, Germany, Austria, Czech Rep., Slovakia, Hungary, Romania, Poland, Ukraine, Moklova, Belarus, Lithuania, Latvia, Estonia, West Russia, Iceland, Greenland (Denmark), United Kingdom, Sweden, Norway, Finland  
  South European: Portugal, Spain, Monaco, Italy, Malta, Slovenia, Croatia, Bosnia and Herzegovina, Serbia and Montenegro, Greece, F.Y.R.O.M (Macedonia), Bulgaria, Turkey
Researcher’s Observations:

**Pigmentation Characteristics**

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<td>Dark Brown □</td>
<td>Light Brown □</td>
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<td>Blue □</td>
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Or describe in your own words:

Notes:
APPENDIX 4 – Published Papers
Hair Color Measurement and Variation

Michelle Vaughn,1* Roland van Oorschot,2 and Swati Baindur-Hudson1

1School of Molecular Sciences, Victoria University, Melbourne, VIC Australia
2Victorian Police Forensic Services Department, Macleod, VIC Australia

KEY WORDS hair pigmentation; reflective spectrophotometry; cluster analysis; forensic science

ABSTRACT Pigmentation of hair in humans has been investigated by medical scientists, anthropologists and, more recently, by forensic scientists. In every investigation, hair color must first be defined by the researchers. Subjective color assessment inhibits the reproducibility of experiments and the direct comparison of results. The aim of this study was to objectively measure human hair color and examine the variation found in a population with European ancestry, using the CIE L*a*b* color space. Observer-perceived hair colors were compared with self-reported hair colors and the color as measured by reflective spectrophotometry of 132 subjects of European ancestry. The presented data show that self-reported hair colors and observer-reported colors are similar; however, these categories are not necessarily the best way to categorize hair color for quantitative research. Using a two-step cluster analysis, hair color can be divided into categories or clusters based on spectrophotometric measurements in the CIE L*a*b* color space and these clusters can be well discriminated from each other. This separation is primarily based on the b* (yellow) color component and the clusters show agreement to observer-reported colors. This study illustrates the possibilities for and necessity of objectively defining the hair color phenotype for various downstream applications. Am J Phys Anthropol 137:91–96, 2008. © 2008 Wiley-Liss, Inc.

Human pigmentation has been observed and studied for over 4000 years (Westerhof, 2006) and is an integral part of our physical health and cultural identity. Normal pigmentation variation in humans has been investigated by medical scientists (Duffy et al., 2004), anthropologists (Parra et al., 2004; Parra, 2007) and, more recently, by forensic science researchers for the prediction of physical characteristics from DNA, to be used as an investigative tool. The prediction of physical traits for use in forensic investigations is a developing field and hair color is one of many traits that may assist investigators in limiting a suspect pool. The UK Forensic Science Service has already implemented a test with 84% accuracy for predicting red hair (Grimes et al., 2001).

Hair color has received somewhat less attention in the study of pigmentation than skin or eye color, with most work on hair color being done as a consequence of, or in conjunction with, other pigmentary traits (such as skin color or melanoma risk) (Sturm et al., 2003; Duffy et al., 2004). These investigations have lead to a great deal of information on red hair (Box et al., 1997; Grimes et al., 2001; Ha et al., 2003; Sturm et al., 2003; Duffy et al., 2004; Naysmith et al., 2004), which is associated with skin cancer risk; however data on non-red hair colors is scarce. Researchers contributing to the scientific investigation of hair color represent the different fields of medical science and genetics (Ancans et al., 2003; Ha et al., 2003; Duffy et al., 2004), the cosmetics industry (Takahashi and Nakamura, 2004, 2005), and forensic science (Grimes et al., 2001), as well as the disciplines of chemistry and toxicology (Borges et al., 2001; Nogueira and Joekes, 2004). There has also been interest in the hair color of our hominid relatives; a recent study has determined that Neanderthals had variants in a gene that is associated with differing pigmentation levels in Homo sapiens (Lalueza-Fox et al., 2007).

The extinct Neanderthals were a hominin species that lived in Europe and western Asia (Krause et al., 2007). Modern Homo sapiens from the same geographical area show almost all of the hair color variation found in human populations. From studies of genetic sequence diversity (Harding et al., 2000), it has been concluded that there was positive selection pressure for maintaining dark skin and hair pigmentation in the southern latitudes of Africa and Asia, where it protected an individual from ultraviolet light from the sun (Rees, 2003; Parra, 2007). When human populations migrated to more northern latitudes, there was less pressure to maintain dark pigmentation and instead, light skin pigmentation became a selective advantage (e.g., through the easier synthesis of Vitamin D) (Rees, 2003; Parra, 2007). The possibility of sexual selection being a factor in the evolution of pigmentation variation is supported by the presence of sexual dimorphism in skin pigmentation and by evidence suggesting that pigmentation influences mate selection (Aoki, 2002; Madrigal and Kelly, 2007a; Parra, 2007); however, difficulties arise in trying to isolate and quantify the effects of natural and sexual selection (Frost, 2007; Madrigal and Kelly, 2007a,b).

Evidence suggests that hair and skin color are somewhat correlated, within the European population (Shriver and Parra, 2000), and hair color is likely the result of similar selective pressures; however, the exact genetic mechanisms of hair color and its high variability, as well as other factors in the biology of hair color have not yet been fully elucidated.

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In every investigation, "hair color" must first be defined by the researchers. What one investigation calls "blonde" however, might be what another calls "light brown" and this inhibits the reproducibility of experiments and the direct comparison of results.

Reflective spectrophotometry has been used to objectively measure skin pigmentation (Shriver and Parra, 2000; Parra et al., 2004) and the technique has also been used on hair in different ways and for various purposes including cosmetic science (Takahashi and Nakamura, 2004, 2005), for the determination of hair composition and structure (Nogueira and Joekes, 2004), for comparison of measurement methods (Shriver and Parra, 2000), and in recent genetic studies (Naysmith et al., 2004). It has been found to be an objective and effective way to measure hair color.

To objectively measure hair color presupposes a method of color measurement. There are, however, numerous models or "color spaces" used to describe color. The model developed by the Commission Internationale de l’Eclairage, CIELAB or CIE L*a*b*, measures color on three axes that are nearly linear with human perception (Ford and Roberts, 1998). This model provides a grid point for each specific color (TASI, 2004) and in addition, has been used for other studies of human pigmentation (Shriver and Parra, 2000; Takahashi and Nakamura, 2004, 2005; Parra, 2007).

In the CIE L*a*b* color space, the lightness, or intensity, of a color is measured on the "L*" axis on a scale from 0 (black) to 100 (white). The color is then measured on the "a*" axis which gives a value from +100 (green) to -100 (red). The "b*" axis measures color from +100 (blue) to +100 (yellow). One unit on the L*, a*, or b* axes is considered to be the smallest difference the human eye can detect (TASI, 2004). This color grid point allows for the mathematical comparison of color. It should be noted that, theoretically, the a and b axes have no maximum or minimum values but this research has used the cut off points of ±100 because these represent the practical limit of the instrument used in the color measurement (Napier S, personal communication, Biolab Group, Australia, 2007).

In 2000, Shriver and Parra used the CIE L*a*b* system with reflective spectrophotometry to measure the hair color of 41 European-American individuals and 18 individuals of non-European ancestry, comparing the L* color component to the Melanin Index. They found a significant correlation between these systems of measurement and reported low variability in the hair color of individuals of non-European ancestry. Naysmith et al. in 2004 used the CIE L*a*b* system to measure the hair color of 50 individuals for genetic and chemical studies of red hair. They found relationships between variations in the gene MC1R and measured hair color (most strongly with b*), as well as with chemical studies. It should be noted that people with red hair were deliberately over-represented in this sample and volunteers ranged in age from 6 to 72 years old (median of 35). Both studies emphasized the necessity and demonstrated the feasibility of measuring pigmentation objectively and accurately.

The aim of the research presented here was to objectively measure human hair color and examine the variation found in a population with European ancestry, over all three color axes. Young adult volunteers were recruited due to the fact that hair color is known to change with age and especially during puberty and late adulthood (Slominski et al., 2004). We were particularly interested in the variation found among the non-red colors (i.e., blonde, brown, and black). Observer-perceived hair colors were compared with self-reported hair colors and the color as measured by reflective spectrophotometry. The variation of the measured hair colors was then analyzed to determine if and how hair colors form describable groups and how these match with our intuitive descriptions of hair color.

Objectively describing categories of hair color will benefit the fields of anthropology and medical science and may introduce some standardization into the design of genetic studies. It could also benefit forensic investigation on various levels, for example, specific criteria may help in better communication between witnesses, investigators, and forensic hair-examiners and be an important part in the future of forensic phenotype-profiling and prediction of physical features from DNA samples.

MATERIALS AND METHODS

Subjects

Subject recruitment and sampling procedures were conducted with the approval of the Victoria University Human Research Ethics Committee. In total, 140 subjects were included in this analysis. All volunteers recruited for the study had their natural hair color at the time of sampling. Most volunteers were between the ages of 18 and 35 and were of European ancestry (132 subjects). The exceptions to these criteria include six individuals of non-European ancestry (one African and five South Asians) and two older European individuals with white hair. Subjects fitting these criteria were chosen to meet the study aims of examining natural variation in adult European hair color, with non-European and mature-white haired individuals included for comparison purposes.

Sampling

Subjects were first given all project information and the following procedures carried out following their written consent (in compliance with Victoria University Human Research and Ethics standards; approval number HRETH06/156). Subjects filled out a questionnaire that, in addition to confirming their age, ancestry, and natural hair color status, asked them to report how they saw their own hair and eye color. Hair color was reported to be one of; Black, Dark Brown, Light Brown, Blonde, or Red. Color assessment by an observer (the researcher taking the samples) was also recorded.

Hair color was measured by reflective spectrophotometry (RS) using a Minolta CR-300 Chroma Meter (Konica Minolta, North Ryde NSW, Australia). The instrument was calibrated using a white tile and a light source input setting of "D65" which represents daylight without spectral highlights. The instrument was set to measure in the CIE L*a*b* format. Hair color was measured on the left, right, and back of the head and was measured five times at each spot, the mean of the 15 measurements being the focus of subsequent statistical considerations. The standard deviation of the 15 measurements for each individual ranged from 0.07 to 4.67 for L*, 0.01 to 2.51 for a*, and 0.03 to 3.3 for b* with means of 1.22, 0.35, and 0.68, respectively.
TABLE 1. The self-reported hair colors (SRC) compared to the observer-reported hair colors (ORC)

<table>
<thead>
<tr>
<th>Color</th>
<th>Self reported</th>
<th>Observer reported</th>
</tr>
</thead>
<tbody>
<tr>
<td>Black</td>
<td>8</td>
<td>12</td>
</tr>
<tr>
<td>Dark Brown</td>
<td>52</td>
<td>44</td>
</tr>
<tr>
<td>Light Brown</td>
<td>37</td>
<td>44</td>
</tr>
<tr>
<td>Blonde</td>
<td>33</td>
<td>31</td>
</tr>
<tr>
<td>Red</td>
<td>8</td>
<td>7</td>
</tr>
<tr>
<td>White</td>
<td>8</td>
<td>12</td>
</tr>
</tbody>
</table>

Statistics

SPSS 15.0 for Windows (©SPSS Inc. 2006, Chicago IL) was used to analyze the color data. Two separate methods were used—a cluster analysis and discriminant analysis. Measured hair color (by RS) was broken into several groups using a two-step cluster analysis; performed using the Euclidean distance criterion or the log-likelihood probability approach and either not specifying the number of clusters to be formed or asking for a specific number of clusters, ranging from two to seven. Cluster analysis seeks to identify natural subgroups within a population by minimizing within-group variation while maximizing between-group variation. To characterize and evaluate the clusters a separate, discriminant analysis was then performed. Discriminant analysis commences from the perspective of there being a known number of population subgroups. From the population, there are a number of individuals whose subgroup classification is known. The data from these individuals is analyzed in an attempt to build a profile of subgroup membership and subsequently use this profile to classify new individuals (whose subgroup membership is otherwise unknown).

After the analysis, SPSS determines which variables are most important in discriminating the groups, and how good this model will be at predicting group membership of future individuals. It determines the predictive value of the model by removing each individual separately, reanalyzing the data, and then predicting the membership of that case. The percent correctly classified is reported.

RESULTS

Reported colors

Self-reported hair colors (SRC) were compared to the colors determined by the observer (observer-reported color or ORC) (Table 1). 85.7% of individuals had the same hair color reported by themselves and the observer. Where there was a difference, discrepancies were lighter or darker by one shade. The observer was more likely (55% of disagreeing observations) to see a darker shade. The observer classified four SRC Dark Browns as Black, four SRC Light Browns as Dark Browns, and three SRC Blondes as Light Brown. The observer also classified eight SRC Dark Browns as Light Browns. In one case, a self-reported Red color was reported as Blonde by the observer.

Population variation

To examine the hair color variation amongst the European population, the L*, a*, and b* values of the 134 European individuals, as measured by reflective spectro-photometry, were recorded and the data subjected to statistical analysis. The variability observed in the entire sample is illustrated in Figure 1. This distinctiveness of the non-European individuals (NE in Fig. 1) and the individuals with white hair (W in Fig. 1) can be seen in the highlighted areas. Individuals not of European ancestry have hair that is much darker with a* and b* values very close to zero.

Grouping the population

The aim of the statistical analysis was to investigate the major components of the color variation and to see whether hair color falls into groups that would be meaningful for further research. What qualifies as a meaningful group is subjective and may depend on the intended use of the data; however in this case, meaningful groups would be those that separate colors into several distinct groups. For practical purposes, it would also be useful to have groupings that correspond well to colors people already use to report hair color (e.g., Blonde).

The European sample was broken into groups using the two-step cluster analysis. The analysis was first conducted without specifying the number of clusters, using either the Euclidean distance approach or the log-likelihood probability approach. Next, the numbers of clusters were specified, from two to seven for both approaches and a discriminant analysis was then performed. Using the clusters determined as a definitive population subgroup, the percentage of cases correctly classified was used to evaluate the clusters.

The percentage of cases correctly classified in the discriminant analysis for the various methods of cluster determination is shown in Figure 2. When the observer-reported color (ORC) groupings are used, only 73.1% of cases are correctly classified. All other clusters have greater than 95% of cases correctly classified.

Determined automatically, without specifying the number of clusters, the Euclidian distance approach...
yielded no useful clusters, while the log-likelihood method grouped the population into two clusters, the composition of which are shown in Table 2 as an example of how clusters are described. When using the Euclidian distance to determine clusters, having two or three clusters results in very poor sample composition, with more than 97% and 93.3% of cases, respectively, in one cluster. From the remaining groups, two and six clusters as determined by the log-likelihood approach (which will be referred to as Analysis 1 and 2, respectively), and four clusters as determined by the Euclidian distance approach (which will be referred to as Analysis 3) showed the highest discriminant scores and the best group composition (data not shown).

During analysis, canonical discriminant functions are determined. These unstandardized functions are used to make classifications in the discriminant analysis and the standardized function coefficients are used to compare the relative importance of each function and each variable in making classifications. The coefficients of the standardized functions shown in Table 3 show that the most informative function in all three analyses is most highly correlated with the \( b^* \) component of color (yellowness) and where there is more than one function, the second most important function is most highly correlated with \( L^* \) (lightness). In Analyses 2 and 3, the third discriminant function contributed 1% and 0.9% of information, respectively (data not shown).

An example on the use of the unstandardized functions is shown in Figure 3, which illustrates the use of the function in discriminating between the two clusters determined in Analysis 1 \( (F = -5.615 + 0.102L^* + 0.212a^* + 0.270b^*) \). Using this function, 97.8% of new cases are correctly classified and this clear distinction between the two groups is visible in Figure 3.

Comparing clusters to the observer-reported colors

It appears that the clusters determined by various methods do show similarity to hair colors as perceived by an observer. When divided into two clusters by the log-likelihood approach (Analysis 1), the observer-reported hair colors Black, Dark Brown, Light Brown, and the four darkest Blondes are in Cluster 1 and the remaining Blondes and all Reds and Whites are in Cluster 2. This shows that the separation in these clusters determined by their \( L^* \), \( a^* \), and \( b^* \) values correspond well to divisions in observer-reported colors and could be referred to as “Dark” and “Fair.” The names are arbitrary; however, they correspond to how people tend to refer to hair color.

The six clusters determined by log-likelihood (Analysis 2) separate the ORC White very clearly and the four Reds with the highest \( a^* \) value (the most red). The other colors are somewhat less distinct; however, they are roughly divided into categories that could be called “Fair,” “Light,” “Medium,” and “Dark.” The example shown in Figure 4 illustrates the separation of the six clusters by applying the two most important discriminant functions (a) and the results of using the same functions on the observer-reported groupings (b), where blurring between the groups can be seen.

When four clusters are determined by the Euclidian Distance approach (Analysis 3), the ORC White is clearly separated, as are the two Reds with the highest \( a^* \) and \( b^* \) values. The rest of the population is divided into two clusters, the first with the darker colors and the other with the fair colors. It should be noted that the separation of fair from dark hair in these groups is similar to that of the two cluster groupings, with four extreme individuals (very light and yellow and the very red) being separated.

**DISCUSSION**

The presented data show that when given a limited choice of colors, self-reported hair colors and those reported by an observer are fairly consistent; however, these categories are not necessarily the best way to cate-
gorize hair color for quantitative research. Although hair color can be perceived as a continuum (as seen in Fig. 1), it can be divided into categories or clusters based on spectrophotometric measurements in the CIE L*a*b* color space and these clusters can be well discriminated from each other. The discrepancies between self-reported hair color and observer-reported hair color and the poor separation of observer-reported color groups compared to those defined by clustering analysis emphasize the importance for objective measurement of individuals to be included in research studies. For repeatability and validity of studies, phenotypes must be defined as well as possible and the objective measurement methods and analysis strategies presented here may be of assistance in this regard.

The discriminant functions used to make classifications show some interesting results. It can be seen that, when using the CIE L*a*b* color space, the b* component (yellow) is the most important in describing the variation and grouping of hair colors, followed closely by L*, in the clusters examined in this study. This suggests that using one component is not sufficient for examining hair colors, as has been done in previous studies (Shriver and Parra, 2000) and that yellow (b*) is the primary contributor instead of L*. This, interestingly, corresponds to the 2004 study by Naysmith et al. (2004) where b* was shown to have the strongest relationship to MC1R genotype. The red component of hair contributes very little information to discriminating groups. The reduction of three color components to one or two discriminant functions makes the examination and classification of individuals easier to represent.

It is also interesting to note that, while not exact, clustered groupings follow the same pattern that self- and observer-reported colors do. How someone decides to report a hair color may be an artifact of the arbitrary naming of colors or of personal bias or it may also be due to human perception of color, which corresponds to the CIE L*a*b* system of measurement. The human eye is a complicated structure with retinal receptors (rods and cones) that are sensitive to changes in lightness and color (Hunt, 1998). Very generally, when receiving a visual signal, these receptors judge the brightness of that signal (mainly the rods) as well as analyzing the hue of a signal as green or red and as blue or yellow (mainly the three types of cones), with this combination of signals being perceived as a color (Hunt, 1998). This system of human trichromatic vision may be why similar groups are defined by both the human reporting of color and by the spectrophotometry and clustering method described here. The lack of sensitivity in biological perception may contribute to the variability found. As mentioned previously, one unit on the L*a*b* scale is defined as the limit of human discrimination between colors (TASI, 2004). The difference between members of different clusters can be less than this in one or more of the three color axes and may contribute to the inconsistency between reported color groups and clusters determined by spectrophotometric data and clustering algorithms.

The particular clustering algorithms and program options used will depend on the intended use of the data. For example, a medical genetic association study may require a Fair/Dark division, where the prediction of hair color in a forensic study may require more specific groupings. This study has illustrated the possibilities by using a two-step cluster analysis in SPSS; however, there are many other programs and clustering algorithms available. Having objectively measured colors allows researchers to group individuals in clearly defined ways and to change these groups as the study demands without subjectivity or collecting additional data.

For practical purposes, it is acknowledged that the exact functions and cluster details may change as larger samples are analyzed and that reflective spectrophotometry, while accurate, may be inconvenient for measurements on very large numbers of people or for investigators and the authors hope to address these issues in the near future.

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**LITERATURE CITED**


Box NF, Wyeth JR, O’Gorman LE, Martin NG, Sturm RA. 1997. Characterization of melanocyte stimulating hormone receptor
A comparison of hair colour measurement by digital image analysis with reflective spectrophotometry

Michelle R. Vaughn a,*, Roland A.H. van Oorschot b, Swati Baindur-Hudson c

Abstract

While reflective spectrophotometry is an established method for measuring macroscopic hair colour, it can be cumbersome to use on a large number of individuals and not all reflective spectrophotometry instruments are easily portable. This study investigates the use of digital photographs to measure hair colour and compares its use to reflective spectrophotometry. An understanding of the accuracy of colour determination by these methods is of relevance when undertaking specific investigations, such as those on the genetics of hair colour. Measurements of hair colour may also be of assistance in cases where a photograph is the only evidence of hair colour available (e.g. surveillance). Using the CIE \(L^* a^* b^*\) colour space, the hair colour of 134 individuals of European ancestry was measured by both reflective spectrophotometry and by digital image analysis (in V++). A moderate correlation was found along all three colour axes, with Pearson correlation coefficients of 0.625, 0.593 and 0.513 for \(L^*\), \(a^*\) and \(b^*\) respectively (\(p\)-values = 0.000), with means being significantly overestimated by digital image analysis for all three colour components (by an average of 33.42, 3.38 and 8.00 for \(L^*\), \(a^*\) and \(b^*\) respectively). When using digital image data to group individuals into clusters previously determined by reflective spectrophotometric analysis using a discriminant analysis, individuals were classified into the correct clusters 85.8% of the time when there were two clusters. The percentage of cases correctly classified decreases as the number of clusters increases. It is concluded that, although more convenient, hair colour measurement from digital images has limited use in situations requiring accurate and consistent measurements.

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Keywords:
Hair colour
Digital image analysis
Reflective spectrophotometry
Forensic science
Genetics
Macroscopic
Investigative tool

1. Introduction

The objective measurement of hair colour has been a topic of several studies in the past few years and is an important part of future research in forensic science [1–4]. Having objectively defined hair colours and the ability to classify them, according to the requirements of a particular study, would allow for increased consistency between research studies and would also allow for greater agreement in discussion between individuals such as investigators, witnesses and forensic scientists. In addition, hair colour is a physical feature that genetic researchers are interested in as part of a ‘phenotype profile’; a way to predict what someone looks like from DNA for investigative purposes [5]. This also requires standardised measurement of hair colour [5]. Reflective spectrophotometry has often been used to measure hair colour and is an established method [1–6]. A previous study by the authors [7] has shown that hair colours measured by reflective spectrophotometry can be classified into meaningful groups, with those groups depending on the intended use of the data. For example, a medical genetics case–control study may be interested in a light/dark separation of colours, where those interested in forensic prediction may be more interested in separating colours into smaller, more specific groups.

While reflective spectrophotometry is an established method for measuring hair colour at the macroscopic level, it can be cumbersome to use on a large number of individuals and not all reflective spectrophotometry instruments are easily portable. This study investigates the use of digital photographs to measure hair colour and compares its use to reflective spectrophotometry. An understanding of the accuracy of the colour determination by these methods is of relevance when undertaking specific investigations, such as those on the genetics of hair colour. Measurement of hair colour by digital image analysis may also be of assistance in cases where a photograph is the only evidence of hair colour available (e.g. surveillance).
Digital image analysis has previously been used to measure hair colour in forensic science; however it has generally been used to measure the colour during the microscopic examination of single hairs and not the macroscopic colour of hair on the head that this study is concerned with [8]. This has been done to assist the traditional forensic hair examination, with the purpose of attributing a hair to a specific donor. Digital images have been used extensively to measure the colour of many other objects, such as vegetables, soil and chocolate [9–11], and have been used to measure skin colour [12–14]. Herbin et al. found that the colour of skin lesions measured from 12 digitised images of colour slides was highly correlated with the smallest difference the human eye can perceive [16]. This grid point allows for the mathematical comparison of colours and also the Red-Green-Blue (RGB) colour space of the computer monitor. Note that digital images are most often displayed and measured in addition to confirming their age, ancestry and natural hair colour status, asked them to report how they saw their own hair colour. Hair colour was reported to be one of; Black, Dark Brown, Light Brown, Blonde, Red or White. Colour assessment by an observer (the researcher taking the samples) was also recorded.

Hair colour was measured by reflective spectrophotometry (RS) using a Minolta CR-300 Chroma Meter (Konica Minolta, North Ryde NSW, Australia). The CR-300 has an 8 mm measurement area and uses D65 viewing geometry. The instrument was calibrated using a white tile and a light source input setting of ‘D65’ which represents daylight without spectral highlights. The instrument was set to measure in the CIE L*a*b* format. Hair colour was measured on the left, right and back of the head and was measured five times at each spot, with the instrument in contact with the head. The mean of the 15 measurements is the focus of subsequent statistical considerations.

Digital images were taken of the left, right and back of the head with a CASIO 4.0 mega pixel camera (model QV-R40) with an image size of 640 × 480 pixels and with picture quality close-up set to ‘fine’. On collection mode (for taking close-up photographs), the flash was turned off, the white balance tool was set to ‘sunlight’ and all other settings were left at the default. Each time a set of images was to be taken under a different light source (on a different day, different time of day, different room, etc.) a piece of new white paper was photographed in the same position as the head of the subject (three sides) and measured using the Minolta CR-300. This ‘white image’ was later used to correct colour for light source. All images were taken at a distance of 15 cm.

2.3. Digital image preparation and analysis

Software packages used included:

- JASC Paint Shop Photo Album 4.0 by Jasc Inc.
- IMAGEJ® 1.0 1994–2006 MathWorks with the Image Processing Toolbox and modules written to convert RGB colour measurements to L*a*b* measurements [9].
- V++ Version 4.0® 1990–2002 Digital Optics Ltd. New Zealand, with modules written to convert RGB colour measurements to L*a*b* measurements [9,18].

Each digital image of the hair was cropped to 100 × 100 pixels in the area closest to the centre of the image (where the RS measurements were taken) with JASC Paint Shop Photo Album, Adobe Photoshop or IrfanView as available. Care was taken to avoid obvious shadows or spectral highlights that result from the curvature of the skull. Initially, images were analysed using MATLAB to convert the RGB colour measurements to L*a*b* and to determine the mean colour values for each image; however, this program is rather difficult and tedious for a new user. The RGB to L*a*b* conversion modules written for MATLAB were instead used within V++ [18], a more user-friendly program, which results in identical values as those determined by MATLAB (results not shown). Therefore, V++ was used to analyse all subsequent images.

For each image, the L*, a* and b* values were calculated. Images of the same colour taken under different light sources will appear different [12], so an effort was made to correct this colour in a manner similar to a previous study using the RGB colour space [12], with the understanding that perfect correction is unlikely and that the greatest difficulty in correcting the colours (a* and b* axes) will be found when the lightness of the image differs. The new white paper, mentioned above, was measured with the Minolta Chroma Meter CR-300 and found to have L*, a* and b* values of 93.4, 1.6 and –2.06 respectively. Using this as a reference point, L*, a* and b* colour points were added or subtracted depending on how much the white reference paper deviated from its Minolta CR-300 determined values under the various sampling situations. For example, if \( L_1 \) is the L* value of the colour image, \( L_2 \) is the L* value of the white image and \( L_3 \) is the L* value of the corrected image, then: \( L_3 = (93.4 – L_1) \times L_2 \).
3. Results

3.1. Colour correction

When comparing the colour mean values of pink paper measured by digital image analysis to the reflective spectrophotometric value, it was found that the DI a* component values have means that are significantly lower than the RS measurements, indicating that the amount of overestimation is not related to the separate colour components (L*, a* and b* axes). A correlation was found along all three axes, with Pearson correlation coefficients of 0.625, 0.593 and 0.513 for L*, a* and b* respectively (p-values = 0.000). This is a significant correlation, but of only moderate strength. The L* axis measurements are most highly correlated, followed by a*, then by b*.

To see if there is a pattern in the differences between DI and RS measurements, a Paired T-test for the difference between the means obtained by DI and RS was performed. Means are significantly overestimated by digital image analysis for all three colour components. The L* value is significantly higher by an average of 33.42 units, the a* value higher by an average of 3.38 units and the b* value higher by an average of 8.00 units (p-values = 0.000).

To investigate whether the overestimation error was uniform across the sample, or if light or dark hair is more prone to overestimation, the difference between the DI and RS measurements was compared to the original hair colour (RS measurement). The differences in L* and b* showed no significant correlation, indicating that the amount of overestimation is not related to the original hair colour for these components; however, there was a low amount of correlation (Pearson’s Coefficient of 0.326) between the overestimation of the DI a* and the RS a* values (p-value = 0.000), indicating that there is a slight tendency for the difference in a* values to become greater as hair becomes more red.

As the separate colour components (L*, a* and b*) measured by RS and DI do not correspond well with each other, we investigated whether when combined they would place individuals into the same colour categories as those determined by RS measurements. In a previous study, we showed that the RS measurements can be used to break this same sample set into defined groups using a Two-Step Cluster analysis [7]. These groups were then evaluated using a discriminant analysis to classify cases into the correct groups. In that study, the best results were obtained for three groups. In this study, the best results were obtained for three analyses: (1) two clusters using a log-likelihood probability method; (2) six clusters using a log-likelihood method and (3) four clusters using a Euclidian distance method. Each analysis obtained classification scores above 95% [7]. This was better than the 73.1% discrimination obtained by using observer-reported clusters [7].

For the present study, each individual was assigned to a cluster based on their RS data (for each of the three analyses). These clusters were then re-evaluated, using their DI data and the percent of cases correctly classified is reported in Table 1. Table 1 indicates that as the number of clusters increases, the percent of individuals correctly classified decreases. Table 1 also shows that
the percent of cases correctly classified in the six observer-reported colour groups (51.5%) was lower than for any of the RS analyses.

Fig. 1 illustrates an example of the use of discriminant functions in classifying individuals (using the results from Analysis 3). The discriminant functions determined by RS data ($F_1 = 0.107 - 0.122L + 0.669a^* + 0.486b^*$ and $F_2 = -8.894 + 0.297L - 0.442a^* - 0.117b^*$) were used to plot individuals onto a grid using first their RS measurements (Fig. 1A) and then their DI measurements (Fig. 1B). Fig. 1A shows the good separation of the four clusters determined by Analysis 3 while Fig. 1B shows the groups merging into one another.

4. Discussion

The presented data have shown that although measuring colour from digital images is convenient, the colours measured and classified do not correspond very well to those colours measured by reflective spectrophotometry and would have limited use in studies requiring accurate measurement of hair colour, such as genetic association studies. They also do not provide an accurate description of hair colour for crime investigators to utilise as visual identifiers or in description announcements. Some use might be found in distinguishing between a small number of clusters with digital images, as in the two clusters of Analysis 1 (which showed 85.8% of cases correctly classified); however, as the number of clusters or groups increases, there is a lower correspondence. Observer-reported colours were not well discriminated by RS data (73.1% of cases correctly classified) [7] and were more poorly discriminated by using digital image analysis (51.5% of cases correctly classified). These results support the assertion that objectively defined and measured colours, rather than observer or self-reported colours are necessary to classify individuals for further study and at this time, this is best achieved through reflective spectrophotometric measurement.

It should be noted that the images in this study were all taken with the same camera under the specific conditions of a well-lit room and from a close, specified and identical distance and all treated in the same way. Multiple analysis of one individual showed very low variation in the results. Any additional variability in conditions would contribute to less similarity between the digital image analysis and a reflective spectrophotometry standard. If variability could be reduced, through improvement in photographic conditions or through more accurate correction for varying conditions, results may improve. The act of photographing an object introduces error in colour reproduction due to the nature of the camera and image production and is an approximation of the visual field (whether digital or with film) [21]. Different models of camera may have different abilities to reproduce colour accurately and may improve results. The process of measuring RGB values and converting to CIE $L^*a^*b^*$ colour space may also result in the loss of some colour information.

It may also be that there are components of hair colour that are omitted when measuring colour directly from a digital photograph. For example, features such as hair texture, thickness, luster or the natural inhomogeneity of hair colours may have a different effect on the analysis of colour from a digital image than they have on reflective spectrophotometry or on human perception. With these things in mind, the improvement of digital cameras and further research into the digital photography of hair would assist in developing methods that provide a more accurate measurement and classification of hair colour that would be of use to researchers, investigators and forensic scientists.

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References


A Comparison of Macroscopic and Microscopic Hair Color Measurements and a Quantification of the Relationship between Hair Color and Thickness

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Abstract: Microscopic images of hair from 36 individuals were evaluated for color (using the CIE \( L^{*}a^{*}b^{*} \) color space), hair thickness, and the pattern of pigmentation. These measurements were compared to each other and to reflective spectrophotometric measurements of macroscopic hair color. Three areas on three strands of hair were examined for each individual. Despite a large \( \Delta E^{*} \) value, there is a strong relationship between macroscopic and microscopic \( L^{*} \) values that would be useful if one was trying to predict the color of a head of hair (macroscopic) from a strand (microscopic). A relationship between hair lightness \( (L^{*}) \) and thickness is also found at both the macroscopic and microscopic levels. The information acquired in this study may be useful to those conducting a microscopic examination of hair, especially in forensic science, as well as those researchers investigating macroscopic hair color and hair morphology for the purposes of forensic science, anthropology, and medical science. Further investigation into objectively measuring and quantifying the microscopic color and properties of hair will assist in increasing its validity as evidence in forensic casework.

Key words: forensic, hair, hair color, microscopic hair, macroscopic hair, hair thickness

INTRODUCTION

Hair is commonly encountered as physical evidence at crime scenes (Saferstein, 2001). Traditional forensic hair examination involves examining the specimen under a microscope and observing characteristics such as color, length, and diameter of the hair, as well as the distribution of pigment granules and other morphological features of the hair that may assist in determining the source of the hair (Saferstein, 2001; Bednarek, 2004; SWGMAT, 2005).

In an attempt to assist with the classification of hair colors in a forensic examination, Bednarek (2004) illustrated one way of measuring hair color at the microscopic level—calculating the red-green-blue (RGB) values of a digital image of the hair. The average RGB values for blond and brown hair were determined. Recent research has also been conducted on the measurement of hair color at the microscopic level for individualizing a hair (Brooks, 2007).

By using a discriminant analysis to assess color measurements, Brooks (2007) was able to correctly assign a strand of hair to a particular individual between 58 and 68% of the time (depending on what color model was used), based on known microscopic color measurements of the individuals’ hair.

While accurate color measurements of hair at the microscopic level are of interest to forensic hair examiners, objectively measured hair color (at both the macroscopic and microscopic level) is also of interest to those investigating the genetic basis of physical features for use in predictive tests (Frudakis, 2008; Vaughn et al., 2008). Reflective spectrophotometry was used to measure the macroscopic hair color in this study because it is an established method and has been shown to be more consistent than digital image analysis (Vaughn et al., 2008, 2009).

The aim of this study was to determine the association between hair color measured at the microscopic level with color measured at the macroscopic level, utilizing the CIE \( L^{*}a^{*}b^{*} \) color space. Because the macroscopic hair color is made up of thousands of strands of slightly different colors, it is important to know if the microscopic color of a strand...
can indicate what the head of hair might look like and vice versa. This would be useful if a strand of hair is found as evidence and possibly also for the prediction of both macroscopic and microscopic color from genetic information. Examining morphological properties of the hair, such as thickness, and their associations with color provides further information on the physical basis of hair color and may also be of relevance to those searching for genetic markers associated with hair color.

**Materials and Methods**

In total, 36 individuals were included in this microscopic analysis. The study included 2 individuals with white hair, 3 individuals of non-European ancestry, and 31 individuals of European ancestry between the ages of 18–35, all with their natural hair color. These subjects were targeted to study natural, adult hair color variation as almost all natural hair color variation is found within individuals of European ancestry (Shriver & Parra, 2000). Each individual had his or her hair color measured by reflective spectrophotometry (RS) as described by Vaughn et al. (2008). In addition, several full-length strands of hair were cut from the scalp of these volunteers for microscopic analysis.

The CIE $L^*a^*b^*$ color space was used for both the macroscopic RS measurements and the microscopic digital image analysis. The CIE $L^*a^*b^*$ model, developed by the Commission Internationale de l’Eclairage, measures color on three axes that correspond to trichromatic human perception and reflect the degree of change in color that humans can perceive (Ford & Roberts, 1998). In this system, the lightness, or intensity, of a color is measured on the $L^*$ axis on a scale from 0 (black) to 100 (white). The color is also measured on the $a^*$ axis that gives a value from $-100$ (green) to $+100$ (red). The $b^*$ axis measures color from $-100$ (blue) and $+100$ (yellow). One unit on the $L^*$, $a^*$, or $b^*$ axes is considered to be the smallest difference the human eye can perceive (TASI, 2004). This grid point allows for the mathematical comparison of colors. It should be noted that, theoretically, the $a^*$ and $b^*$ axes have no maximum or minimum values. Our research has used the cut off point of $\pm 100$ because this is the practical limit of the software and instrument used to measure color (Napier, 2007).

Hairs were mounted on Livingstone Premium glass slides (76.2 × 25.4 mm pathology grade) with Entellan mounting medium (Merck) and covered with Marienfeld cover slips (22 × 60 mm No. 1). The area to be examined was marked on the slide. Three random areas, at least 1 cm away from the hair root or tip, on each of three hairs were examined for each individual, and the mean of these measurements was used for most analyses (Fig. 1).

**Equipment and Software Used to Generate Microscopic Images and to Measure Color**

Equipment and methods are from Brooks (2007) with minor modification.

Leica DMLB is a microscope with a 50× oil immersion lens (500× magnification; field of 276 μm), using 10.5 V illumination (daylight equivalent) and with DLF (daylight filter) on.

Olympus DP70 is a 12.5 megapixel digital camera (Olympus Optical Co. Ltd. 2003), and the Image Management System consists of the DP Controller (Version 2.1.1.183) and DP Manager (Version 2.1.1.163). An image size of $1360 \times 1024$ pixels was used with ISO 200 sensitivity, exposure time set to “automatic,” 1% (of the field) contrast correction, and all images saved as TIF files.

Automontage-Pro 5.02.0096 software (by Syncroscopy, a division of Synoptics Group, UK) was used to montage images of the middle third of the hair (the area containing the greatest amount of pigmentation information) (Brooks, 2007). Hair thickness was estimated by the total depth of the optical field (Fig. 2).

V++ Version 4.0 (by Digital Optics Ltd. New Zealand, ©1990–2002) with a module written to convert RGB color measurements to $L^*a^*b^*$ measurements was used to extract
the correct areas of the microscopic images and to measure the color of the extracted area (Figs. 3, 4).

**RESULTS**

**Comparing Microscopic Color to Macroscopic Color**

A difference in color can be measured by calculating \( \Delta E^* \), an Euclidian measure of the distance between two colors in the CIE \( L^*a^*b^* \) space (Hunt, 1998):

\[
\Delta E^* = \left[ (\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2 \right]^{1/2}.
\]

The average \( \Delta E^* \) between the macroscopic and microscopic color was 22.57 units, with a standard deviation of 5.41 units. However, this measurement does not give information about which components are different, and therefore each color axis was examined separately.

When comparing microscopic (Mc) color \( L^* \), \( a^* \), and \( b^* \) measurements directly to the macroscopic measurements determined by RS, regression analysis found that \( a^* \) and \( b^* \) measurements fail to show any significant linear relationship between Mc and RS (\( p \)-values = 0.317 and 0.288, respectively). RS and Mc \( L^* \) show a moderate, significant linear relationship with an \( R = 0.631 \) (\( R^2 = 0.469, p \)-value = 0.000). However, using curve estimation (multiple regression), it can be seen that there is a stronger quadratic relationship between the variables (Fig. 5).

**Comparing Macroscopic Color to Hair Thickness**

Macroscopic hair color, measured by RS, was compared to the estimates of hair thickness (the two white-haired individuals were not included). Macroscopic \( L^* \) (Fig. 6) and \( b^* \) were found to have a moderate, negative correlation with hair thickness, with correlation coefficients of \(-0.648 \) and \(-0.657 \) (\( p \)-values = 0.000), respectively. The \( a^* \) value showed a significant (\( p \)-value = 0.023) but weaker correlation of \(-0.390 \). This means that thinner hair is correlated with being lighter and more yellow.

![Figure 3. Extracted image of a light brown hair.](image)

![Figure 4. Extracted image of a blond hair.](image)

![Figure 5. The quadratic relationship between microscopic \( L^* \) measurements and \( L^* \) measurements by RS. The trendline has the equation \( y = 0.0188x^2 - 1.2854x + 48.503 \) and \( R^2 = 0.8164 \).](image)
Comparing Microscopic Colors to Hair Thickness

Microscopic hair color was compared to the estimates of hair thickness. There were significant, negative correlations between the microscopic $L^*$ values of hairs and their estimated thickness (in micrometers) with an $R = -0.705$ ($p$-value = 0.000) (Fig. 7). The $a^*$ value was positively correlated to the hair thickness ($R = 0.567$, $p$-value 0.000), while $b^*$ showed no significant relationship (data not shown).

DISCUSSION

This study illustrates the relationships between macroscopic and microscopic color and hair thickness. It illustrates that, despite the large $\Delta E^*$ value, there is a strong relationship between macroscopic and microscopic $L^*$ values that would be useful if one were trying to predict the color of a head of hair (macroscopic) from a strand (microscopic) or if one...
was trying to predict hair color phenotype by genetic analysis and either the macroscopic or microscopic hair color was known to be more closely associated with a particular genetic variation. The relationship between hair lightness ($L^*$) and thickness is also clearly illustrated at both the macroscopic and microscopic levels.

While the reported relationships are useful and will assist in further research toward the objective measurement of hair colors, it can also been seen that there are weak or insignificant relationships between some aspects of the macroscopic and microscopic color, particularly when examining the $a^*$ and $b^*$ color components. These differences may be due to the fact that the number of hair strands that can be examined under the microscope were, and in the course of normal casework investigation most likely will be, a small proportion of the head of hair and larger sample sizes are difficult to obtain.

Color measurement by digital image analysis also presents some problems. In a previous study (Vaughn et al., 2009), the authors have shown that macroscopic hair color measured by digital images analysis does not correspond very well to the RS readings of the microscopic hair color, and some of the same factors may be involved in measuring the color of microscopic images, such as the ability of the camera to reproduce color and the conversion from RGB to the $L^*a^*b^*$ color space. However, the high degree of consistency in conditions under a microscope should make microscopic digital images of hairs comparable to each other and therefore more consistent in their relationships with the macroscopic color.

The measurement of hair thickness could also be improved to increase the strength of relationships between thickness and color. Cross sections of the hair may be examined to determine the shape before measuring the thickness of hairs because using the optical depth may inadvertently use different sides of an oval-shaped hair for measurement and therefore be less precise.

The information acquired in this study may be useful to those conducting the microscopic examination of hair, especially in forensic science, as well as those researchers investigating macroscopic hair color and hair morphology for the purposes of forensic science, anthropology, and medical science. Further investigation into objectively measuring and quantifying the microscopic color and properties of hair will assist in increasing its validity as evidence in forensic casework.

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