INTRINSIC FACTORS AFFECTING DECOMPOSITION
CHANGES IN ARCHAEOLOGICAL HEAD HAIR
FROM KELLIS 2 CEMETERY, DAKHLEH OASIS, EGYPT

by

KATHLEEN COLE
B.A. University of Central Florida, 2012

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ABSTRACT

Post-mortem hair root degradation, and associated characteristics such as post-mortem root banding, brush-like ends, and hard keratin points, has remained a little understood phenomenon in the forensics discipline since its discovery in the 1800’s. At present, the underlying causes of these characteristics are still unknown. In addition, there is no standardization for preparing samples for forensic or archaeological analysis. In this study, 1200 hairs from a total of 51 individuals (males, n = 22; females, n = 29) ranging in age from 16 to 60+ and interred at the Kellis 2 cemetery in the Dakhleh Oasis, Egypt are examined microscopically for evidence of post-mortem hair root degradation. These remains date from ~50AD to ~450AD, and all were naturally mummified. The purposes of this thesis are two-fold; 1) to determine the efficacy of two preparation methods, and 2) to examine the intrinsic and extrinsic variables in each hair in order to ascertain the factors that affect the degradation of the human hair root after death.

The preparation methods include a dry sample, where the hair is directly removed from the scalp tissue using tweezers, and a wet sample, where a 1cm portion of the scalp was first rehydrated using dimethyl sulfoxide (DMSO) for 48 hours before the hairs were extracted. Results indicate that when working with naturally mummified remains the wet method provides for easier acquisition of the sample and less chance of breakage before the hair is mounted for observation. This, therefore, provides a larger sample size for analysis.

Each hair was observed using polarizing microscopy to determine whether postmortem root degradation was present, the growth stage of the hair, color, pigment density and distribution, pigment aggregate size and shape, medulla continuity and
opacity, cuticle scale profile and thickness, inner cuticle margin, and the presence of ovoid bodies. Significant correlations were found between the incidence of postmortem root degradation and the growth stage, hair color, cuticle thickness, cuticle scale profile, and individual's age. The remaining variables did not show any significant correlations.
In Loving Memory of Granddad, Nano, Aunt Kathy, and Granddad Jack

You are deeply missed.
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LIST OF ACRONYMS/ABBREVIATIONS

AP- Arrector Pili Muscle
BLE- Brush-Like Ends
CSP- Cuticle Scale Profile
DMSO- Dimethyl Sulfoxide
DOP- Dakhleh Oasis Project
DP- Dermal Papilla
HKP- Hard Keratin Point
IRS- Inner Root Sheath
ORS- Outer Root Sheath
PHRM- Postmortem Hair Root Morphology
PMRB- Postmortem Root Banding
CHAPTER 1: INTRODUCTION

Hair analysis techniques were first entered into forensic evidence in the 1800’s, and since then numerous case studies have examined the usefulness of hair trace evidence in medico-legal cases (e.g., Gaudette et al. 1985; Petraco et al. 1988; Linch and Prahalow 2001; Collier 2005; Koch et al. 2013). Detailed studies have been conducted to determine the evidential significance of hair in DNA analysis, drug and toxicological analysis, and elemental analysis (e.g., Gordus 1973; Ishiyama et al. 1982 Wilson et al. 1995; Uhl 1997; Tagliaro et al. 1998; Budowle et al. 2003; Kintz 2004; Goullé et al. 2004; Pragst and Balikova 2006; Pillay and Kuis 2007). Additionally, comparative analyses have been utilized to determine the possible sources of questioned hairs in cases where the origin is unknown (e.g., Miller 1987; Smith and Goodman 1995-1996; Deedrick and Koch 2004; Bisbing 2007). In situations such as these, hairs and fibers are collected from known sources and compared to the questioned hair with the hopes that a match can be made and the source of the questioned hair fiber discovered. Despite the strides made by previous researchers and investigators regarding forensic hair analysis, minimal research has been conducted examining the morphological changes that occur to hair, specifically to the proximal root-end of human hair, during the process of decomposition, regardless of its potential use as evidence in forensic investigations (Linch and Prahalow 2001:15).

Currently, these hair fiber root-end changes are classified using the terms postmortem root banding (PMRB), hard keratin points (HKP), and brush-like ends (BLE). PMRB refers to “an opaque ellipsoidal band which appears to be composed of a collection of parallel elongated air spaces [that] is approximately 0.5 mm above the root bulb and
about 2 mm below the skin surface” usually assumed to occur in the stages of early decomposition (Petraco et al. 1988:73). BLE and HKP, on the other hand, are usually thought to be associated with late term decomposition. In these cases, “the hair will break off at the area of the dark band often leaving the proximal end discolored and with a pointed or brush like appearance” (Koch et al. 2013: S52).

When these changes were first discovered by Japanese scientists Hajime Sato and S. Seta in 1984, they referred to these phenomena collectively as 'putrid roots', a term which is still in limited use today (Ort 2005; Kadane 2015). Following their discovery, the Committee on Forensic Hair Comparison (1985), “in an effort to advance forensic hair comparison as a science,” called for a standardization of the terminology to be used by hair analysts, and the term “putrid root” was added to the list (Gaudette et al. 1985:1, 8). These phenomena were not studied in depth until 1988 when Petraco et al. (1988) wrote “The Morphology and Evidential Significance of Human Hair Roots”, the first in a very short list of research conducted in an attempt to understand the causal factors associated with the morphological changes observed on human hair roots after death. Unfortunately, this study only discussed two types of putrid roots in depth, PHRB and BLE. The third type, HKP, was only briefly mentioned. This is likely due to the probability that PMRB, BLE, and HKP are related to different types and stages of decomposition, and the research conducted by Petraco et al. (1988) was limited in both scope and scale. A study examining all of the postmortem changes to human hair would have required a large and diverse sample set, which was not available at the time. According to Koch et al. (2013) the need for a large study sample might explain the paucity of research which has been conducted on PHRM to date – as samples from cadavers are not easily acquired (S52).
It is without question that researchers face many difficulties pertaining to studies involving deceased individuals, but it is imperative that these phenomena undergo rigorous research. Since many investigations of PHRM (e.g., Petraco et al. 1988; Linch and Prahlow 2001; Collier 2005; Koch et al. 2013) have found a link between changes of the human hair root and decomposition of the body itself, its detection and association with an active crime scene can potentially indicate that a deceased individual was present even in cases where a body is missing. In short, hair trace evidence holds great potential for medico-legal investigations, especially where postmortem changes in the hair root may be the only indication that a crime has been committed.

**Medico-legal Use of Postmortem Changes in the Hair Root as Evidence**

In 2005, *People v. Kogut*, 10 Misc. 3d 245, 247, 805, N.Y.S.2d 789, 791 (N.Y. Sup. Ct. 2005) set the precedent for admission of PMRB evidence in the 2011 case of the *State of Florida v. Casey Marie Anthony* case number: 48-2008-CF-015606-O (2011a:3-5). *People v. Kogut*, 10 Misc. 3d 245, 247, 805, N.Y.S.2d 789, 791 (N.Y. Sup. Ct. 2005), however, was a retrial of a murder case which took place in 1986 with Kogut and two others convicted of the murder of Theresa Fusco. In the original trial, Nicholas Petraco, who is a leading forensic microscopist and trace evidence analyst, compared two questioned hairs found in a van belonging to one of the defendants to known hairs that were collected from the victim. He stated that the questioned hairs were found to be consistent with the known hairs, thereby suggesting they came from the same individual (Ort 2005:1). Despite having observed PMRB on the roots in 1986, this trait was not discussed until the 2005 retrial of *People v. Kogut*, 10 Misc. 3d 245, 247, 805, N.Y.S.2d 789, 791 (N.Y. Sup. Ct. 2005).
Petraco, who testified for the prosecution in the 1986 trial, was now testifying for the defense along with Peter De Forest, a forensic expert who was also a witness for the defense in 1986. Both De Forest and Petraco testified that PMRB "occurs only when the hair begins to decompose inside the hair follicle, while still on the head of the deceased" and that "the victim's body would not have been in the van long enough for postmortem hair banding to occur" (Ort 2005:1). Since the hair from the vehicle was eerily similar to the hairs taken from the body during autopsy, and since those hairs were stored in unsealed enveloped, their testimony led to the belief that that the evidence had been contaminated and commingling had occurred between hair samples (Innocence Project 2015). After 19 years of incarceration, Kogut was finally found innocent, and in 2012 litigation began on a wrongful conviction suit where, again, PMRB is likely going to play an important role (Seybert 2012: 1-36).

Another case where postmortem root banding was introduced as evidence was the high-profile trial of the State of Florida v. Casey Marie Anthony case number: 48-2008-CF-015606-O. This case began in July 2008 when Caylee Anthony, Casey Anthony’s only child, was reported missing by her grandmother, Cindy, who stated to a 911 dispatcher that, “I have a possible missing child. I have a three-year-old that’s been missing for a month” (State of Florida v. Casey Marie Anthony, 48-2008-CF-015606-O 2011b; Montaldo 2015). Later in the day, during a subsequent call to 911, she told the dispatcher, “I found my daughter’s car today and it smells like there’s been a dead body in the damn car” (Battaglia 2011/2012:1587; State of Florida v. Casey Marie Anthony, 48-2008-CF-015606-O 2011b; Montaldo 2015). As it turned out, Caylee had last been seen alive on June 16th of that year, but her mother had never reported her missing. She told her mother and
police that she had been searching for Caylee on her own, and that her daughter had been abducted by her nanny. Changes in her original story, and the accompanying circumstances, resulted in her being named as the number one suspect in the disappearance of her daughter; she was arrested on July 16th, 2008. Unfortunately, despite many searches of the property and surrounding woods, no body was found. Casey Anthony was later released after posting bond, but searches of the car, residence, and computer search history led police to the conclusion that Caylee was likely deceased. On October 14th, 2008 Casey was rearrested and charged with murder, manslaughter, and lying to police, all of which took place without the presence of Caylee’s remains.

On December 11th, 2008, the skeletonized remains of a child were discovered in the woods adjacent to the Anthony’s neighborhood. The remains were confirmed as those of Caylee Anthony on December 19th, 2008. Evidence of possible homicide was immediately apparent as duct tape was adhered to the skull where the mouth would have been located, and also due to the method and placement of body during disposal. The real test for investigators was linking Casey Anthony to the death of her child. Hair collected from the trunk of the car became part of the State’s evidence that Caylee’s body had been placed into the trunk, and that Casey had indeed killed her own daughter.

Steven Shaw, an FBI hair examiner, testified at trial that the hair acquired from the left side of the trunk liner was consistent with hairs obtained from the trash bag containing Caylee’s remains. However, he noted that, “hairs are not a means of positive identification” stating that he could not say “a hair did originate from a person to the exclusion of all others” (State of Florida v. Casey Marie Anthony, 48-2008-CF-015606-O 2011b). Additionally, mtDNA testing was conducted on exhibit Q12.1 (a human head hair)
and compared to DNA obtained from a buccal sample of Casey, but since mtDNA is passed directly from mother to child, Dr. Catherine Theisen, who provided the analysis, stated that “neither Casey Anthony nor Caylee Marie Anthony can be excluded as the source of the Q12.1 hair” (Theisen 2008:2). Since prosecutors could not use DNA or hair comparison to exclude Casey as the source of the Q12.1 hair, they relied on evidence of postmortem root banding in order to show that the hair belonged to an individual who was deceased, and therefore, could not have belonged to Casey or Cindy Anthony.

Karen Korsberg Lowe, a trace evidence analyst for the FBI laboratories, examined the Q12.1 hair and noted possible evidence of decomposition in her report. She stated that “a Caucasian hair found in specimen Q12 exhibits characteristics of apparent decomposition at the proximal (root) end”, but no other hairs were found in the evidence collected that showed signs of putrid root formation (Theisen 2008:2). When called to the stand to testify, Lowe stated that based upon her experience working with trace evidence and on the work conducted by previous researchers, postmortem root banding was found to be consistent with decomposition. Further, she did not know of any studies where PMRB, like that observed on Q12.1, had been replicated in hairs acquired from a living person (State of Florida v. Casey Marie Anthony, 48-2008-CF-015606-O 2011a). Despite her expert testimony and previous experience, she admitted that there were only a handful of studies conducted on PMRB. While this evidence may not have decided the case for Casey Anthony, it is possible that a greater amount of scientific knowledge could have buttressed the forensic hair analysis leading to a far different conclusion. On July 5th, 2011, the jury found Casey Anthony not guilty for the murder of her daughter. On July 17th, 2011, three years after Caylee went missing, Casey was released from jail.
Another case involved the murder of two adult males, a mother, and her son. In this case, hair was analyzed from the crime scene and compared to known samples acquired from the remains of each individual (Tafaro 2000). The hairs collected from the two adult male victims were microscopically differentiated from the mother and her son, but the hairs from the mother and her son could not be excluded from one another. When microscopically examined it was found that the juvenile’s hair contained the only evidence of PMRB, while his mother’s hair showed only BLE. In this way, investigators were able to establish which hairs belonged to the mother and which belonged to her son. These observations further helped to corroborate the suspect’s confession that he had returned several days after the murder to slit the throats of all the victims. This also helped to establish how hairs showing decomposition changes were found on knives discovered at the crime scene (Tafaro 2000:497-498).

**Research Focus**

This thesis includes an examination of postmortem hair root morphology (PHRM) utilizing hair samples obtained from the scalps of individuals buried in the Kellis 2 cemetery in the Dakhleh Oasis, Egypt. It has been suggested by previous researchers that the presence of postmortem changes in the proximal root-end of the human hair, with an emphasis being placed specifically on PHRB, could be a tell-tale sign that a deceased individual was present at a location even if a body cannot be found. Therefore, the impact this type of evidence can have on a medico-legal investigation is tremendous, and yet our understanding of PHRM is suffering from a serious lack of scholarly research.

Fortunately, there are a few studies which discuss extrinsic factors that affect PHRM, but literature discussing intrinsic factors- traits specific to the hair itself and the
individual- is nonexistent. Therefore, in an effort to fully understand the many factors which may have an effect on PHRM, it is necessary to study the intrinsic variables as well. This is mainly because individuals tend to die under various sets of circumstances and experience different taphonomic environments. Given that no two hairs are exactly the same, it is crucial to determine why postmortem changes occur in some hair roots, while not occurring in others even within a single individual’s remains.

While most researchers suggest that PMRB, HKP, and BLE occur only after death, the scarcity of studies related to these phenomena and the circumstances surrounding their appearance leads to questions and complications when hair samples demonstrating these traits are admitted into evidence. Therefore, at the focus of this thesis is the hair root morphology of samples procured from the naturally mummified remains of individuals interred at the Kellis 2 cemetery, Dakhleh Oasis, Egypt, in order to determine which variables affect the appearance of postmortem changes in human hair roots. These samples were archived by the Dakhleh Oasis Project (DOP) and housed at the University of Central Florida Laboratory for Bioarchaeological Research. The hairs included in this study were collected from individuals interred within the burials highlighted in red in Figure 1.1. Additionally, utilizing hair samples provided by the DOP will ensure a study sample of known provenience and relative environmental stability. It is hoped that this research will add to the scientific and forensic knowledge surrounding PHRM, aid and promote future research within the field, and thereby provide a solid foundation for the use of PHRM as evidence in future legal proceedings.
Figure 1.1: Map of the Kellis 2 cemetery. Burials of individuals included in this study are highlighted in red. (Adapted from Williams 2008)
The Dakhleh Oasis Project

The Dakhleh Oasis is located approximately 660Km SSW of Cairo in Egypt's Western Desert (Dupras 1999). It is one of five major depressions and is known for its hyper-arid environment. Temperatures vary from -4°C to 25°C in winter and from 19°C to 50°C during the summer months (Blume et al. 1984, Giddy 1987, Dupras 1999, Wheeler 2010). Rainfall is also rare, with 0.3 mm/year being the average (Blume et al. 1984). High atmospheric pressure, violent winds (Khamasins), and seasonal sandstorms further contribute to the aridity in the oasis (Dupras 1999). It is believed that the climate has remained relatively stable, with very little change, from the time that ancient Kellis was inhabited, and the cemeteries were in use, until now. Because of this, the level of preservation for many of the remains is remarkable. Additionally, since the remains at Kellis 2 were naturally, rather than artificially, mummified, tissue samples, such as skin, internal organs, muscle tissue, hair, and nails, were suitable for this study (Williams 2008). Figure 1.2 shows the approximate extent of the Dakhleh Oasis and its geographical location in relationship to Cairo, Egypt.

The excavations that have taken place at the Kellis 2 cemetery constitute a significant portion of the research focus of the Dakhleh Oasis Project. This project involves the excavation and subsequent analyses of archaeological sites within the Dakhleh Oasis, Egypt (Bagnall 1997; Cook 1994). The Dakhleh Oasis Project (DOP) began in 1978, under the direction of Anthony Mills, and has continued into the present with the support of various universities and organizations; among them The Royal Ontario Museum and the Society for the Study of Egyptian Antiquities (Dupras 1999). The bioarchaeologists involved in the Dakhleh Oasis Project have excavated numerous
inhumations at the west and east cemeteries (Kellis 1 [K1] and Kellis 2 [K2] respectively) associated with the ancient town of Kellis (Birrell 1999). Of importance for purposes of this thesis, are the excavations at Kellis 2 which began in 1992 (Birrell 1999) and have continued to the present day.

Figure 1.2: Map showing the approximate extent of the Dakhleh Oasis and its geographical relationship in relationship to Cairo, Egypt. (Google Earth)
As of 2012, a total of 771 burials have been excavated from this site (701 have been analyzed), but it is estimated that the Kellis 2 cemetery contains between 3,000-4,000 graves in total (Molto 2002; Wheeler 2010).

Kellis 2 dates from around 50 AD to about 450 AD, the Roman-Byzantine period in Egypt, and is assumed to be an early Christian cemetery based on grave and body orientation, and the mortuary practices observed at the site (Stewart et al. 2003; Molto 2002). The burials were generally single interments, where the body was wrapped in a linen shroud, placed in an extended, supine position within a mud brick tomb, and situated with the head to the west and the feet to the east; very few graves included a coffin of any kind and many were lacking grave goods (Birrell 1999; Williams 2008; Wheeler 2009). Additionally, the burials of infants and children were situated alongside those of adults, demonstrating that this population believed in the humanness of all individuals within their community. This is an idea that was lacking in Egypt during its earlier periods, but which is consistent with later Christian ideals (Bowen 2003; Bowen et al. 2005). The period during which this cemetery was in use also coincides with the appearance of the first Christians in Egypt during the first two centuries AD (Bowen 2003; Wheeler 2010). Additionally, the remains have been subjected to radiocarbon dating that has tentatively dated Kellis 2 to the late third and early fourth centuries AD, suggesting that the cemetery may have been in use for longer than was originally assumed (Birrell 1999; Molto 2001; Stewart et al. 2003).

Research Questions

Three research questions will be addressed in this thesis; 1) What is the best method of sample preparation for the observation of PHRM in a mummified sample?; 2)
Which intrinsic factors affect whether decomposition changes affect the hair root?; and, 3) Is postmortem root banding indicative of early stage decomposition? It is necessary to understand which factors are related to the absence or presence of putrid roots. As such, each hair in this study will be evaluated for extrinsic factors such as environmental degradation, microbial action, and insect activity, as well as intrinsic attributes including: growth stage, medulla characteristics, pigment characteristics, color, presence or absence of ovoid bodies, and cuticle characteristics in order to analyze the factors related to the appearance of PHRB, HKP, and BLE. Additionally, biological sex, age, and health status (where available) of the individuals will also be noted to provide a more comprehensive overview of the study sample. The overarching goal of this study is to establish a standard methodology for other researchers wishing to study PHRM.

**Thesis Structure**

This thesis is structured in an integrated article format. Every chapter is set up in such a way as to facilitate dissemination in peer-reviewed publications.

Chapter 2: The Examination of Two Preparation Methods for the Study of Postmortem Hair Root Morphology, introduces two methodologies employed in this study. These two methods are utilized to determine which method is most beneficial when working with dry decomposition. An overview of the processes for preparing wet and dry samples, for mounting of the samples on to slides for observation, and an analysis of the efficacy of each method in relation to the study sample are presented.

Chapter 3: Intrinsic Factors Affecting Postmortem Hair Root Morphology, focuses on intrinsic factors and how they affect the occurrence of PHRM. The variables included are age, biological sex, growth stage, hair color, pigment density, pigment distribution,
pigment aggregate size, pigment aggregate shape, medulla continuity, medulla opacity, cuticle thickness, inner cuticle margin, cuticle scale profile, and ovoid bodies. This chapter includes archetypes for each of these variables, as well as a discussion of the statistical analyses performed and the variables’ relationships to PHRM.

Finally, Chapter 4 concludes this thesis with an overview of the results and offers suggestions for applications and further research regarding PHRM.
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Wilson, Mark R., Joseph A. DiZinno, Deborah Polanskey, Jeri Replogle, Bruce Budowle.
CHAPTER 2: THE EXAMINATION OF TWO PREPARATION METHODS FOR THE STUDY OF POST-MORTEM HAIR ROOT MORPHOLOGY

Forensic hair examination is not new to the field of forensic science. An 1857 study by John Glaster, *Hairs of Mammalia from the Medico-Legal Aspect* (1931; cf. Evans 2004:101-102), discusses the significance of hair from a medico-legal standpoint and is still regarded as a reliable reference on the subject. Additionally, the first evidential use of human hair in an American legal case occurred during June of 1882 in the state of Wisconsin (Ellis 1882). *Knoll v. State* was a landmark murder case in that it was the first to include the examination of human hair in witness testimony. This being a new field of forensic science, however, the examination consisted of a single hair comparison between the crime scene and victim. In addition to the minute amount of evidence, the examination of the hair itself was conducted with the naked eye rather than microscopically; a fact that had the judge questioning its validity. In truth, the most the ‘expert witness’, Dr. Piper, could offer was that “the hair was precisely the same in every respect, in length, magnitude, color, and in every other respect” (Ellis 1882:381). Even with this paltry evidence, he concluded, “as a result of that comparison, I can say that it was from the head of the same person” (Ellis 1882:381).

Forensic hair analysis has come quite a long way from this starting point to include toxicology, DNA, microscopic hair and fiber comparisons, and postmortem hair root morphology (PHRM). Unlike many forensic hair applications, which have established specific methodologies for analyses, PHRM is a relatively new area, and as such, standards have yet to be established. Therefore, the purpose of this research is to
determine which preparation methodology will provide the most consistent and usable results when examining PHRM for use as evidence in medico-legal cases. This study specifically looks at whether preparation method affects the retention of a usable sample, the decomposition changes observed, and the growth stages obtained. Growth stages include anagen or active growth stage, catagen or transitional growth stage, and telogen or terminal growth stage (Ogle and Fox 1999). In this case, growth stages are of particular importance since it has already been established through previous research that growth stages have an effect on whether decomposition changes are observed or not. A greater proportion of anagen and catagen hairs within the sample, hairs that have been shown to demonstrate postmortem changes in the hair root, should yield substantially different results than a sample consisting mainly of telogen or terminal growth stage hairs. Since telogen hairs have ceased growth and the root has become fully keratinized, postmortem changes are no longer observed (Petraco et al. 1988; Linch and Prahlow 2001; Koch et al. 2013). Therefore, a method that provides a greater proportion of anagen and catagen hairs would be significantly more effective.

It is hoped this research will contribute to the establishment of methodological protocols for future forensic analysts. The hair used in this particular study originated from the remains of individuals associated with the Kellis 2 cemetery (~50AD-~450AD) in the Dakhleh Oasis, Egypt, and was acquired by bioarchaeologists involved in the Dakhleh Oasis Project (DOP).

Dakhleh Oasis Project (DOP)

The Dakhleh Oasis is one of the seven main Oases of Egypt (Figure 2.1). It is located approximately 660Km SSW of Cairo and 300Km west of Luxor in Egypt’s Western
Desert (Dupras 1999). Dakhleh Oasis boasts a hyper-arid environment, where temperatures can range anywhere from -4°C in the winter months to 50°C during the summer (Blume et al. 1984; Giddy 1987; Dupras 1999; Wheeler 2010). Annually, only 0.3mm of rainfall is accumulated (Blume et al. 1984). According to Dupras and Schwarcz (2001), “humidity can range from 23% to 30% from March to September, rising to 33% to 50% from October to February” (1200). *Khamasins*, which are violent winds that ravage this area, sandstorms, and high atmospheric pressure, serve to increase aridity in this already arid environment (Dupras 1999).

Archaeological excavations of the Kellis 2 (K2) cemetery in the Dakhleh Oasis have been ongoing since 1991 (Cook 1994:260). The K2 cemetery dates to circa AD 50 to about AD 450, which coincides with the Roman-Byzantine period and the appearance of the first Christians in Egypt. This is also consistent with radiocarbon dating conducted on the remains, which tentatively dates the cemetery to the late third to early fourth centuries AD and suggests that the cemetery may have been in use for longer than previously assumed (Birrell 1999; Molto 2001; Stewart et al. 2003). Additionally, the mortuary practices observed at Kellis 2 are consistent with Christian mortuary practices from the same era (Stewart et al. 2003; Molto 2002). These burials consisted mostly of single interments with the deceased buried with their head facing towards the west and their feet towards the east; the graves themselves contained little to no grave goods (Birrell 1999; Williams 2008; Wheeler 2009). The inclusion of children and infants into the cemetery populations lends further credence to the assumption that this is an early Christian cemetery since children were not included in previous cemetery populations (Bowen 2003; Bowen et al. 2005).
To date, 701 of the 770 individuals excavated have been examined and analyzed by DOP bioarchaeology team members. Because the environment in Dakhleh Oasis has remained consistent over the past 2000 years, the human remains from the Kellis 2 cemetery are very well preserved. Many of the individuals buried in the Kellis 2 cemetery underwent natural, or spontaneous, mummification, allowing for the preservation of tissues such as hair.

Figure 2.1: Map showing the approximate extent of the Dakhleh Oasis and its geographical relationship in relationship to Cairo, Egypt. (Google Earth)
Materials and Methods

For this study, a total of 400 individual hairs from 10 individuals from the Kellis 2 cemetery were analyzed, and were used for the purpose of determining the best method for sample preparation. The scalp samples are currently stored in the Laboratory for Bioarchaeology Research at the University of Central Florida (UCF). Both males and females are included in the study population and their ages range from between 16-21 up to 60+. Tables 2.1-2.3 provides a breakdown of the sample IDs included in the study population, age categories for each individual, and the proportion of males to females. Additionally, Figure 2.2 shows the extent of the excavations conducted at Kellis 2 cemetery as well as the distribution of those individuals that have been included in this research.

Table 2.1: Sample IDs, age, biological sex, and number of hairs for each individual included in the study population.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Age</th>
<th>Biological Sex</th>
<th># of Individual Hairs Acquired</th>
</tr>
</thead>
<tbody>
<tr>
<td>B261</td>
<td>60+</td>
<td>F</td>
<td>40</td>
</tr>
<tr>
<td>B265</td>
<td>36-50</td>
<td>M</td>
<td>40</td>
</tr>
<tr>
<td>B269</td>
<td>51-60</td>
<td>F</td>
<td>40</td>
</tr>
<tr>
<td>B279</td>
<td>22-35</td>
<td>F</td>
<td>40</td>
</tr>
<tr>
<td>B280</td>
<td>60+</td>
<td>F</td>
<td>40</td>
</tr>
<tr>
<td>B281</td>
<td>60+</td>
<td>M</td>
<td>40</td>
</tr>
<tr>
<td>B284</td>
<td>22-35</td>
<td>F</td>
<td>40</td>
</tr>
<tr>
<td>B291</td>
<td>36-50</td>
<td>F</td>
<td>40</td>
</tr>
<tr>
<td>B306</td>
<td>60+</td>
<td>F</td>
<td>40</td>
</tr>
<tr>
<td>B309</td>
<td>22-35</td>
<td>M</td>
<td>40</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td>400</td>
</tr>
</tbody>
</table>
### Table 2.2: Age categories and distribution for individuals included in the study population

<table>
<thead>
<tr>
<th>Age Range</th>
<th># of Individuals in Age Category</th>
<th># of Hairs Included in Sample Population</th>
<th>% of Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>22-35</td>
<td>3</td>
<td>120</td>
<td>30.0</td>
</tr>
<tr>
<td>36-50</td>
<td>2</td>
<td>80</td>
<td>20.0</td>
</tr>
<tr>
<td>51-60</td>
<td>1</td>
<td>40</td>
<td>10.0</td>
</tr>
<tr>
<td>60+</td>
<td>4</td>
<td>160</td>
<td>40.0</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>10</strong></td>
<td><strong>400</strong></td>
<td><strong>100.0</strong></td>
</tr>
</tbody>
</table>

### Table 2.3: Distribution of biological sex for the study population

<table>
<thead>
<tr>
<th>Sex</th>
<th>Number</th>
<th>% Total Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>F</td>
<td>280</td>
<td>70.0</td>
</tr>
<tr>
<td>M</td>
<td>120</td>
<td>30.0</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>400</strong></td>
<td><strong>100.0</strong></td>
</tr>
</tbody>
</table>
Figure 2.2: The Kellis 2 cemetery excavations conducted through 2007. Red graves are those individuals whose remains were used in this study (Adapted from Williams 2008)
Sample Preparation

Forty hairs with roots attached were acquired from each individual using two different methods. For the first preparation method, referred to as the dry method, ten individuals were chosen based upon the significant amount of hair and scalp that had been collected in the field. Age and sex data were provided by Dr. Lana Williams based on previous research conducted by the DOP (Williams, 2008). Each hair was individually extracted directly from the scalp, examined to confirm that the root was attached, and placed in labeled bags. Without any further processing, the first 20 hairs were bagged and labeled with the individual’s identification number.

The second preparation method, referred to as the wet method, utilized the same ten individuals included in the first method to ensure that there was no prejudice in the sample based on an individual’s characteristics or level of decomposition. A square measuring 1cm x 1cm was cut from each scalp section and placed in a separate petri dish where it was rehydrated using dimethyl sulfoxide (DMSO) for 24-48 hours following the method outlined by Williams (2008). Twenty hairs were extracted from each of these samples, allowed to dry overnight, and then placed in labeled bags. In this method, DMSO was used to rehydrate the scalp to make it easier to pluck the individual hairs and lessen any possible damage the root. This method was also used to determine whether the preparation would affect the occurrence of decomposition changes in the hair root.

Five slides were prepared and labeled for each individual and the preparation method was noted on each slide. Permount® was chosen as a mounting medium due to its refractive index (RI) of 1.525 at 25°C being remarkably close to the RI of human hair at ~1.55 (Linch and Prahlow 2001; Marschner et al. 2003; Committee on Identifying the
Needs of the Forensic Sciences Community 2009:156). Four hairs were mounted on each slide using Permount®, creating five slides for the dry method and five slides for the wet method for each individual. All slides were left to dry for 24-48 hours before being stored to ensure that the mounting medium was fully set. The specimens were then examined using an Amscope 40x-1500x Infinity Polarizing Microscope at 40x and 60x magnification. Data was collected on the presence or absence of hair roots (i.e., were the roots lost after they were bagged or during the sample preparation), growth stages (e.g., were more hairs in an active growth stage present in one method versus the other), differences in the rate of postmortem root banding (PMRB), brush-like ends (BLE), and hard keratin points (HKP).

The data was input into SPSS® statistics software for statistical analysis to determine which method provided the most definitive results. Chi-square analysis was conducted to ascertain whether methodology affects the number of usable samples obtained from the remains, the growth stages present within the study sample, the incidence of postmortem changes seen, and the type of postmortem changes observed.

Results

The results show that in cases where the body and tissues succumb to dry decomposition, it is beneficial to use the wet method for procurement of intact hair and hair root samples. In fact, out of 200 hairs included in the dry method, 36% (72/200) of the hair samples had no roots, whereas using the wet method only about 20% (39/200) of the hairs sampled lost their roots during processing (Table 2.4).
Table 2.4: Cross-tabulation analysis of presence and absence of roots using two dry and wet preparation methods.

<table>
<thead>
<tr>
<th>Method</th>
<th>No Roots</th>
<th>Roots</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry</td>
<td>72</td>
<td>128</td>
<td>200</td>
</tr>
<tr>
<td>Wet</td>
<td>39</td>
<td>161</td>
<td>200</td>
</tr>
<tr>
<td>Total</td>
<td>111</td>
<td>289</td>
<td>400</td>
</tr>
</tbody>
</table>

The growth stage for each sample was noted, except in cases where the root was no longer present, to determine if the method of preparation had an effect on which growth stages were observed in each sample. The results indicate that the preparation method did affect the observed growth stages. In the wet method, more anagen and catagen roots were present, while in the dry method the main roots that were intact were those in the telogen growth stage (Table 2.5). Results from the Chi-square analysis indicate that the correlation between the method of preparation and the growth stages observed is significant at the p=0.01 level (Table 2.6). This is more than likely due to how each of the hair samples were procured for each method. For example, anagen and catagen roots were more easily obtained intact from the rehydrated scalp sample, while in the dry samples the majority of intact roots were in the telogen growth stage. This is because telogen roots tend to survive the process of pulling the hair out of the dry scalp better than anagen and catagen roots, which are softer and more easily damaged during the collection process and which retain anatomical structures affixing them to the dermal layer of the scalp.
Table 2.5: Incidence of each growth stage observed using each preparation method

<table>
<thead>
<tr>
<th>Growth Stage</th>
<th>Method</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Dry</td>
<td>Wet</td>
</tr>
<tr>
<td>Anagen</td>
<td>28</td>
<td>93</td>
</tr>
<tr>
<td>Catagen</td>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td>Telogen</td>
<td>78</td>
<td>55</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>113</strong></td>
<td><strong>152</strong></td>
</tr>
</tbody>
</table>

Table 2.6: Chi-Square analysis for preparation method and growth stage

<table>
<thead>
<tr>
<th></th>
<th>Value</th>
<th>Asymp. Sig. (2-sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pearson Chi-Square</td>
<td>51.024a</td>
<td>.000</td>
</tr>
<tr>
<td>Likelihood Ratio</td>
<td>53.130</td>
<td>.000</td>
</tr>
<tr>
<td>N of Valid Cases</td>
<td>400</td>
<td></td>
</tr>
</tbody>
</table>

a. 0 cells (0.0%) have expected count less than 5. The minimum expected count is 5.50.

Lastly, to determine which preparation method yielded the most definitive results, the occurrence of PMRB, BLE, and HKP was documented. Since, as noted above, the preparation method affected the growth stages procured, it is not surprising that the decomposition changes observed also differed. The decomposition changes most observed with the dry preparation method were both PMRB and BLE; together these made up about 10% of the total sample (11/200 PMRB and 11/200 BLE). In the wet samples, however, PMRB accounted for over 25% (53/200) of the total sample, and only five BLEs were observed (Table 2.7).
Table 2.7: Frequencies of postmortem changes in conjunction with preparation method

<table>
<thead>
<tr>
<th>Decomposition Stage</th>
<th>Method</th>
<th>Dry</th>
<th>Wet</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMRB</td>
<td>11</td>
<td>53</td>
<td>64</td>
<td></td>
</tr>
<tr>
<td>BLE</td>
<td>11</td>
<td>0</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>BLE &amp; PMRB</td>
<td>0</td>
<td>5</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>HKP</td>
<td>3</td>
<td>0</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>N/A</td>
<td>103</td>
<td>103</td>
<td>206</td>
<td></td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>128</strong></td>
<td><strong>161</strong></td>
<td><strong>289</strong></td>
<td></td>
</tr>
</tbody>
</table>

The analyses suggest that when dry decomposition is encountered, the most fitting method to use, to acquire intact roots directly from the scalp of the deceased, is likely the wet method. Not only did this method lessen the amount of damage observed on the roots, but it can also be suggested that when BLE are encountered it could be the result of hair that was forcibly removed from an already desiccated set of remains.

**Discussion**

The analyses suggest that when dry decomposition is encountered, the most fitting method to use, to acquire intact roots directly from the scalp of the deceased, is likely the wet method. Furthermore, it could be stated that the wet method would be beneficial regardless of the state or type of decomposition observed in the remains. An individual’s body tends to lose moisture as it decomposes making it more difficult to obtain intact and usable hair samples if the scalp is not properly rehydrated. A drier scalp could lead to a disproportionate number of hairs in the telogen growth stage or hairs showing brush-like ends or hard keratin points.
Since telogen or terminal growth stage hairs are resistant to postmortem change, acquiring a sample that includes a greater proportion of hairs in the anagen and catagen growth stages would be preferable. Undamaged anagen and catagen roots are more easily obtained from a rehydrated scalp, meaning the wet method is the method that should be followed. Not only did the wet method provide a more usable sample for this research (i.e. there was a greater proportion of anagen and catagen hairs in the wet sample than were in the dry sample, 97:35 respectively), but it can also be suggested that when BLE and HKP were encountered in the study sample it may be the result of the hair being forcibly removed from an already desiccated set of remains. The fact that PMRB was observed five times as much in the wet sample as the dry sample (53:11 respectively) and there were no BLE observed in the wet sample further supports this assumption.

**Conclusion**

This study examined two sample preparation methods for use by forensic investigators when obtaining intact hair root samples from mummified remains. The main purpose of this research was to test two preparation methods to determine which is better for preserving postmortem hair characteristics. The results of this research, examining dry and wet methods, indicates that the wet method provided more hairs in the active growth stage (anagen) and transitional growth stage (catagen). Additionally, the forceful plucking of hair in the dry method caused the loss of many of the hair roots both during the process of obtaining the hair and during the time when the hair was stored before being mounted.
The majority of the hairs with lost roots were most likely in the anagen and catagen growth stages since most of the roots that remained intact were in telogen stage, the final hair stage where the hair has ceased growth. Since decomposition changes were not observed in telogen roots, the dry method did not provide as much material as the wet method in regards to studying postmortem human hair root morphological changes. Additionally, since telogen hairs do not appear to show evidence of decomposition, it is even more integral to a forensic investigation to obtain intact hairs and hair roots in the anagen and catagen growth stages. As such, the wet method appears to be the most useful and advantageous method of preparation for forensic investigators trying to obtain hair samples from mummified remains.
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Molto, J.E. 

Ogle, Jr Robert R., Michelle J Fox.  

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Stewart, J.D., J.E. Molto, P.J. Reimer.  
Wheeler, Sandra M.

Wheeler, Sandra M.

Williams, Lana J.
CHAPTER 3: INTRINSIC FACTORS AFFECTING POSTMORTEM HAIR ROOT MORPHOLOGY

A number of studies have examined the effect that extrinsic factors have on the appearance of postmortem hair root morphology (PHRM), but very few researchers have sought to determine whether intrinsic factors affect these changes. Factors such as age, biological sex, pigment, medulla, and cuticle characteristics can significantly affect the type and/or extent of postmortem changes observed. Unfortunately, these have not been analyzed by previous researchers. The only intrinsic growth factor that has been studied in depth is hair growth stages.

Previous research (e.g. Petraco et al. 1988; Linch and Prahlow 2001; Koch et al. 2013) has shown that hairs in the anagen or active growth stage and those in the catagen or transitional growth stage are the only hairs subject to postmortem hair root degradation. Hairs in telogen or the terminal growth stage do not show decomposition changes. It is believed that this is due to the full keratinization of the telogen hair root. Since anagen and catagen stage hairs are not fully keratinized, they are more susceptible to decomposition changes. Given that growth stages have been found to adversely affect whether postmortem hair root degradation is observed, it is not a stretch to believe that PHRM might be affected by other intrinsic factors such as those listed above. Therefore, the purpose of this study is to examine the possible links between the occurrence of postmortem root banding (PMRB), brush-like ends (BLE), and hard keratin points (HKP), and the intrinsic characteristics that may affect PHRM. A total of 713 hair samples, acquired from the scalp sections of 51 individuals interred at the Kellis 2 cemetery (Figure 3.1) in the Dakhleh Oasis, Egypt, composed the study sample.
Figure 3.1: Map of the Kellis 2 cemetery. Burials of individuals included in this study are highlighted in red. (Adapted from Williams 2008)
The Dakhleh Oasis is located in Egypt's Western Desert and approximately 660Km SSW of Cairo (Dupras 1999). The Oasis is known for its hyper-arid environment and temperatures there can range from -4°C in the winter months to 50°C during the summer (Blume et al. 1984, Giddy 1987, Dupras 1999, Wheeler 2010). Rainfall is rare, with an average annual accumulation of only 0.3 mm (Blume et al. 1984). According to Dupras and Schwarcz (2001), “humidity can range from 23% to 30% from March to September, rising to 33% to 50% from October to February” (1200). Violent winds known as *Khamasins*, sandstorms, and high atmospheric pressure also contribute to this hyper-arid environment (Dupras 1999). Because the environment in Dakhleh Oasis has remained relatively stable from the time Kellis was inhabited to the present day, the level of preservation, despite the absence of artificial mummification procedures, is phenomenal. Additionally, since the remains at Kellis 2 were naturally, rather than artificially, mummified, samples from these individuals are more easily acquired (Williams 2008). Figure 3.2 shows the approximate extent of the Dakhleh Oasis and its geographical location in relationship to Cairo, Egypt.
Excavations have been ongoing at the Dakhleh Oasis since the project began in 1978, with those at Kellis 2 beginning in 1991 (Birrell 1999). The K2 cemetery itself dates from around 50 AD to about 450 AD. This time frame is contemporaneous with the Roman-Byzantine period and the appearance of the first Christians in Egypt. The Kellis 2 cemetery is an early Christian cemetery based on the mortuary practices observed at the
site as well as body positioning (Stewart et al. 2003; Molto 2002). The burials were generally single interments, except in a few cases where two bodies (mother and child or twins) were found buried together. Additionally, bodies were usually wrapped in linen shrouds and placed in an extended, supine position. The body was situated with the head facing to the west and the feet to the east; very few graves included any type of coffin or grave goods (Birrell 1999; Williams 2008; Wheeler 2009). Additionally, the inclusion of infants and children into the general cemetery population is strictly associated with Christian ideology, which touts the humanness of all individuals (Bowen 2003; Bowen et al. 2005). Some of these remains have also been radiocarbon dated, resulting in a tentative date of the late third to early fourth centuries AD. This suggests that the cemetery may have been in use for a longer period of time than was originally thought (Birrell 1999; Molto 2001; Stewart et al. 2003).

**Background and Significance**

Very few studies have been conducted with the sole purpose of examining PHRM. In fact, Linch and Prahlow (2001) state that “there are... no references for detailed descriptions of hair root decomposition changes” (15). This, however, is not surprising. Even though hair analysis has been part of the field of forensics for some time, decomposition changes of the human hair root (also termed putrid roots) were not discussed until 1984 by S. Seta and Hajime Sato (Kadane 2015). Later, Petraco et al. (1988) added to the subject in the article *Morphology and Evidential Significance of Human Hair Roots*. Theirs was the first known study conducted specifically with the purpose of observing postmortem human hair root morphology and discussing what factors affect its appearance.
Koch et al. (2013) suggest that the main reason for the dearth of research conducted in this area is because of the need for a large number of cadavers that can be placed under controlled circumstances and regularly monitored (S52). Controlling the circumstances under which a body decomposes is indeed a problem since environmental conditions have a direct impact on the decomposition changes observed; a very strict protocol would need to be followed to ensure the accuracy of the study.

Fortunately, while studies regarding intrinsic hair characteristics are sorely lacking, research has been conducted to determine whether environmental and other extrinsic factors affect the occurrence and appearance of Postmortem Hair Root Morphology (PHRM). For instance, Lynch and Prahlow (2001) state that according to their experience, “the brush like hair proximal end appeared to more commonly arise from dry scalp area, while the hair keratin point hair ends appeared to more commonly arise from moist scalp areas” (16). The environment, as well as body positioning, can also affect the occurrence of PHRM. For example, an exposed scalp can yield a drier mode of decomposition than a scalp that is directly in contact with the ground. This is also dependent on the mode of deposition. Bodies disposed of in water will undergo very different decomposition changes than those that are placed in a hyper-arid environment, and these differences extend to the postmortem changes observable in the human hair root.

Fortunately, even though the K2 cemetery contains individuals buried over many centuries and who died almost two thousand years ago, this environment offers a research advantage. The majority of the remains were buried in a similar fashion, and with similar grave goods. While some minor environmental fluctuations may have occurred, the climate has been relatively stable since the time of burial up until the
present. In addition, none of the bodies underwent artificial mummification. Not only does this mean that the bodies decomposed naturally, it also means that the decomposition changes should be mostly consistent with little variation throughout the study sample. This being the case, it can be assumed that all of the hair samples used in this research should yield similar results if only affected by extrinsic factors.

**Morphological Changes Occurring to Human Hair during Decomposition**

Specific terminology is currently used to describe changes that occur to the hair proximal end and root after death and during the process of decomposition. These terms are postmortem root banding (PMRB), hard keratin points (HKP), and brush-like ends (BLE). PMRB is usually described as “an opaque ellipsoidal band which appears to be composed of a collection of parallel elongated air spaces [that] is approximately 0.5 mm above the root bulb and about 2 mm below the skin surface” (Petraco et al. 1988:73). Figure 3.3 provides a good example of PMRB, which is generally thought by forensic hair analysts to be associated with the earlier stages of decomposition. As decomposition advances, “the hair will break off at the area of the dark band often leaving the proximal end discolored and with a pointed or brush like appearance” (Koch et al. 2013: S52). While this does not always occur, when HKP and BLE are present it suggests that the body of the deceased is in an advanced stage of decomposition even if the body is not present for examination. Examples of BLE and HKP are provided in Figure 3.3 for comparison.
(a) Postmortem Root Banding

(b) Brush-like ends

(c) Hard keratin point

Figure 3.3: (a) Postmortem Root Banding (PMRB) observable on the proximal end of the human hair. PMRB is generally considered to be associated with the early stages of decomposition. This hair was acquired from K2 B306. (b) This is typical of brush-like ends, which are generally thought to result from the later stages of dry decomposition. This hair was acquired from K2 B204. (c) Typically thought to result from the later stages of moist decomposition, Hard Keratin Points were rare within this sample. This example was acquired from K2 B309.
Although there is, without a doubt, a lack of material written on the subject of putrid roots, there have been a few researchers who have studied PHRM. For instance, Linch and Prahlow’s (2001) discuss the decomposition criteria and extrinsic factors thought to be associated with the occurrence of these morphological changes. They found that while BLE were generally associated with dry decomposition, HKP resulted from moist decomposition (Linch and Prahlow 2001:18). Additionally, PMRB seemed to occur early on in the decomposition process, later being replaced by BLE or HKP as decomposition advanced. Koch et al. (2013) further refined the categories of PMRB, HKP, and BLE. Based on their observation of human hair roots, they presented a scoring system for any of the observed morphological changes. A score of 0 meant that no changes were observed, while a score of 1 indicated the beginning stages of root banding. A score of 2 was an indication that a full band was observed, and finally a score of 3 meant that the root had broken off leaving a brush-like end; archetypes were provided for each of these scores (Koch et al. 2013: S54-S55).

While more studies are needed on the subject of postmortem morphological changes that may occur to human hair, several publications (Petraco et al. 1988:74; Linch and Prahlow 2001:16; Jeffery et al. 2007:289; Koch et al. 2013: S58) agree that PMRB, BLE, and HKP seem to appear exclusively in anagen or late anagen/early catagen roots. Because of the keratinization process that occurs during the telogen growth stage, telogen hairs appear to remain mostly unaffected by decomposition changes. The keratinization process causes hairs in the telogen stage to be nearly impervious to alteration (Domzalski 2004:2; Jeffery et al. 2007:290). Researchers (e.g., Petraco et al. 1988; Linch and Prahlow 2001; Koch et al. 2013) also seem to agree that PMRB occurs
during the earlier stages of decomposition, while HKP or BLE appear to occur as decomposition progresses. According to most researchers, their appearance is also conditional upon the type of decomposition (wet versus dry) associated with the remains (Petraco et al. 1988:75; Linch and Prahlow 2001:19; Jeffery et al. 2007:290; Koch et al. 2013:S52). It should be noted, however, that it is not uncommon to see more than one type of hair root decomposition change occurring within a single set of remains. This can depend on body positioning, environmental circumstances surrounding the individual's death, and the asynchronous pattern with which human hair grows during an individual's life (Koch et al. 2013:S58). Asynchronous hair growth means that human hair does not grow in a predictable, seasonal pattern, unlike the hair growth cycles seen in other mammals.

Inarguably, one of the most notable points that researchers seem to agree upon is that putrid roots, in their many forms, are only observed after death. Though there have been studies which have attempted to recreate these morphological characteristics in hair acquired from living individuals (Domzalski 2004), none appear to have been completely successful (though see Linch and Prahlow 2001:19). In fact, it seems that no researchers up to the present have successfully pinpointed why these changes occur in some situations, but fail to appear in others. Again, this is not because there is a lack of interest in the field, nor because it is of little importance to the forensic science community, but rather due to the need for a large number of cadavers that can be placed under controlled circumstances and regularly monitored (Koch et al. 2013:S52). Due to the limitations placed on this type of research, very few attempts have been made to study, *en masse*, the various factors that might affect the appearance of putrid roots in any one individual.
Fortunately, even though there are still very few published articles on the subject of putrid roots, some researchers have made it a point to study the effects that environmental factors, climate, microbial action, time since death, and burial environment have on human hair. For instance, in court records for *People v. Kogut*, 10 Misc. 3d 245, 247, 805, N.Y.S.2d 789, 791 (N.Y. Sup. Ct. 2005), Seybert (2012) cited independent studies conducted by De Forest, Collins, and Houck who studied the effects of various environmental factors on PHRM. According to their testimony, De Forest stated that he believed “microbial action may be to blame for PMRB”, while Collins’ research suggested that “PMRB development is correlated with temperature” (as cited in Seybert 2012:13, 27). Other case studies have focused on time since death and its effect on PHRM; although, notably, none can agree on how much time has to pass after death before putrid roots appear. Researchers such as Houck (as cited in Seybert 2012), assert that the hair must be “in the decomposing skin for a minimum of 7 days”, but this time frame is not agreed upon by everyone in the forensic science community (Petraco et al. 1988:74; Seybert 2012:9).

Unfortunately, unlike extrinsic factors that have received some interest in forensic literature, intrinsic characteristics, those qualities relating to the individual involved and the hair itself, have received little to no attention in these studies. The intrinsic factor that has garnered the most attention, and the only one that appears to have been established by previous researchers (e.g., Petraco et al. 1998; Linch and Prahlow 2001; Koch et al. 2013) as affecting hair after death, is the hair’s growth stage. The growth stage is a major factor since PMRB and other degenerative changes only occur on anagen and early catagen roots. Since these roots are not shed naturally, their presence at a crime scene
suggests that a struggle likely took place, with the anagen and catagen stage hairs being forcibly removed from the scalp. In conjunction with the appearance of PMRB, HKP, or BLE it further suggests that decomposition has possibly occurred, and that it is no longer simply a case of assault. Despite the usefulness of such research, growth stage appears to be the only intrinsic factor actively studied by previous researchers (e.g., Petraco et al. 1998; Linch and Prahwol 2001; Koch et al. 2013); other qualities, both macroscopic and microscopic, have not accumulated as much interest.

Since human hair possesses a myriad of traits that make each hair unique unto itself, it is just as imperative to study these in order to determine which of them might affect PHRM. As outlined in several publications, these traits include: hair form (straight, curved, wavy, loose curl, tight curl); color (light, light to medium, medium to dark, and dark); pigment qualities (density, distribution, granule size, and aggregate shape and size); and medulla opacify and continuity (Gaudette et al. 1985:52, 54-56; Bencze 1994:207-208; Ogle and Fox 1999:20-40; Deedrick and Koch 2004). Studying any one of a combination of these traits might help to determine why some individuals show evidence of PMRB, HKP, and BLE, while others do not.

Hair Structure and Biology

In order to discuss the intrinsic variables included in this study, it is important to understand the structure and biology of hair itself. For most mammals, hair develops in utero. This is because “hair is beneficial as insulation from the very moment of birth” (Morioka 2005:4). For humans, hair development begins at the eighth week gestation period and continues until birth. At approximately eight weeks after conception, a cluster of cells forms a bulge under the epidermis. This is known as the pre-germ stage, and it is
soon followed by the hair-germ stage at the eleventh week gestation period. During this stage, the dermal papilla, small protrusions of the dermis, begin to form and melanocytes, pigment cells that provide the hair color, can be discerned in the epidermis. It is also at this stage that the hair follicle will begin to take on the slanted position obtained once it has been fully formed (Harding and Rogers 1999:2-3). The hair-germ stage is followed by the hair-peg stage and then the bulbous-peg stage. Finally, at around the 19th to 21st week gestation period the hair follicle is fully formed, and the fetus will be covered in a fine, downy layer of lanugo (fetal hairs) or vellus hairs (non-medullated hairs) that will be consistently shed and replaced until birth, although, this can continue into the post-natal period (Wollina 1997:33; Harding and Rogers 1999:4; Sinclair et al. 1999:3).

**Intrinsic Factors**

The hair follicle can be divided into four zones; “(1) the cell proliferation and differentiation zone at the base of the bulb, (2) the keratin gene expression zone in the upper bulb, (3) the keratogenous zone in which the hardening of the fiber occurs, and (4) the zone of IRS [inner root sheath] degradation” (Harding and Rogers 1999:7). These zones are also known as the bulb, the suprabulbar area, the isthmus, and the infundibulum. The bulb is comprised of the dermal papilla (DP) and epithelial matrix cells, while the suprabulbar area and isthmus are areas in which the inner root sheath of the hair begins and ends. Finally, the infundibulum includes the sebaceous gland and the arrector pili muscle (AP) (Harding and Rogers 1999:5-6; Torkamani et al. 2014:88). The sebaceous gland is responsible for producing a waxy material called sebum, which helps to waterproof the hair and lubricate the skin of mammals. The AP, on the other hand, “mediates thermoregulation by contracting to increase air-trapping, but was thought to be
vestigial in humans” (Torkamani et al. 2014:88). For most mammals, when the AP is contracted it allows the hair to stand on end providing extra insulation or a useful intimidation tactic by making the animal appear larger and more formidable. This is not the case with humans given how unnoticeable human hair appears. An overview of the hair follicle and surrounding tissues can be seen in Figure 3.4.

Figure 3.4: Diagram of human hair follicle, hair shaft, and surrounding tissues.

Figure 3.5 provides a closer look of the proximal end of the hair shaft itself, as well as its various parts. From medial to lateral, the hair bulb or root consists of the dermal papilla (DP), the basement membrane surrounding the DP, the medulla, cortex, and the cuticle. The hair fiber is then encased in the inner root sheath (IRS) consisting of the IRS cuticle layer, Huxley layer, and Henle layer, and this is surrounded by the outer root sheath (ORS). The ORS is “the most external component, [which] encases the other cell layers
of the follicle and is contiguous with the epidermis” (Harding and Rogers 1999:5). The following sections of this chapter provide an overview for each part of the hair shaft examined in this thesis. Also provided are the archetypes for each variable included in the study sample.

![Diagram of the inner structure of the human hair shaft](image)

**Figure 3.5:** Diagram illustrating the inner structure of the human hair shaft.

**Growth Stages**

Humans undergo intermittent hair growth, where “each hair grows to a maximum length, is retained for a period of time without further growth and is eventually shed and replaced” (Sinclair et al. 1999:7-9); this is a constant process known as the hair growth cycle. The term cycle, itself, is problematic because “it implies a periodic regularity and
this is not the case in human hair growth” (Harding and Rogers 1999:40). Given the regular use of this terminology in scientific journals and publication, the term cycle is used in this study to refer to the three growth stages which hair undergoes.

Hair begins this cycle in the anagen growth stage. During this hair is actively growing. This length of time that this stage lasts depends mostly on where the hair is found on the body. For vellus and arm hairs, the anagen stage lasts for a short period of time (about 6-12 weeks). However, leg hair can remain in the anagen stage for 19-26 weeks, and scalp hair can stay in anagen stage for 2-5 years (Sinclair et al. 1999:9; Jeffery et al. 2007:282). On average, 80-90% of human hair is in the anagen (active growth) stage (Petraco et al. 1988:69; Robertson 1999:86; Koch et al. 2013:S53). According to Robertson (1999) “the average human has in the order of 100,000 hairs”, which means that about 80,000-90,000 of these are actively growing (86). A good example of a hair root during anagen (active growth stage) can be seen below in example (a) for Figure 3.6.

Following anagen stage is a period of regression or transition; this is known as the catagen stage. Catagen hairs account for only ~1-2% of hair on the human scalp (Petraco et al. 1988:69; Koch et al. 2013: S53). This is because the catagen stage of hair growth lasts for only 2-3 weeks (Jeffery et al. 2007:282). During the catagen stage, the dermal papilla and lower portion of the hair follicle retract upwards, and the hair root begins to take on the club-like appearance it will obtain during the final stage (Whiting and Dy 2008:55). Given the low frequency of hairs in the catagen stage, there were very few examples in the study sample. Additionally, the catagen stage is arguably the most difficult growth stage to determine morphologically. Unlike the anagen and telogen stages, which
are characterized by their distinctive root morphology, the catagen stage lies somewhere between anagen and telogen in appearance, making identification of this stage challenging. Despite their rarity, however, there were a few examples found within the study sample. A hair root in catagen is shown in Figure 3.6b below.

The hair cycle finishes when the hair root reaches telogen stage. This stage accounts for the final ~10-20% of hair in the scalp (Petraco et al. 1988:69; Koch et al. 2013:S53). Hair generally spends a total of 2-4 months in telogen stage before finally falling out (Harding and Rogers 1999:44; Jeffery et al. 2007:282). During telogen, the root takes on a club-like appearance, and is visible to the naked eye due to its lack of pigment (Harding and Roger 1999:45; Robbins 2002:9). Telogen hairs constitute the greatest portion of hairs found on clothing and in forensic cases since they are usually naturally shed, when a new hair pushes the telogen hair out, or brushed out. In fact, “compared with hairs in anagen stage, they require only a small force to dislodge them” (Harding and Roger 1999:44-45). A hair root in telogen can be seen below in Figure 3.6c.
(a) Typical appearance for the human hair root during the anagen growth stage. This hair was acquired from K2 B289.

(b) Typical appearance for the human hair root during the catagen growth stage. This hair was acquired from K2 B131.

(c) Typical appearance for the human hair root during the telogen growth stage. This hair was acquired from K2 B242.

Figure 3.6 Human hair growth stages archetype reference chart for Kellis 2 sample population.
Hair Color

Follicular melanocytes, similar to epidermal melanocytes, are responsible for giving each individual hair its unique color. The color itself is generally the product of various amounts of melanin contained within the hair follicle, a process that is ultimately determined through genetics. “Melanin granules are mainly in the cortex, their long axis parallel to the hair surface”, and they are generally a mix of two types of melanin; eumelans and pheomelans (Castanet and Ortonne 1997:213). These are not exclusive, but rather both types can be present within a single hair follicle. Eumelans are associated with black to brown pigmentation and are usually associated with darker hair, while lighter hair results from pheomelans that are reddish yellow (Harding and Rogers 1999:21; Robbins 2002:179; Joblonski 2004:590).

The Atlas of Human Hair (1999) provides a list of seven hair colors and the label of “other” for colors that do not fit into these categories. For instance, hair that has been dyed blue would be placed into the “other” category since it is not a color appearing naturally in human hair. The colors provided by the Atlas of Human Hair (Ogle and Fox 1999) were used as archetypes for the hairs examined in this study. Following the classification structure provided in the Atlas of Human Hair (Ogle and Fox 1999), Table 3.1 provides a list of the hair colors included in the study sample utilized in this thesis, while Figure 3.7 shows some of the archetypes as they appeared within the study sample. As can be seen from Table 3.1, the majority of the study sample included darker hair colors such as dark golden brown. However, blonde and red hair were also present, as were grey and colorless (white) hair.
<table>
<thead>
<tr>
<th>Color</th>
<th>Category</th>
<th>Frequency in Sample</th>
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</thead>
<tbody>
<tr>
<td>Colorless</td>
<td></td>
<td>130</td>
</tr>
<tr>
<td>Blond</td>
<td>Light</td>
<td>66</td>
</tr>
<tr>
<td></td>
<td>Medium</td>
<td>24</td>
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<tr>
<td></td>
<td>Dark</td>
<td>8</td>
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<tr>
<td>Golden Brown</td>
<td>Light</td>
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<tr>
<td></td>
<td>Dark</td>
<td>145</td>
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<tr>
<td>Brown</td>
<td>Light</td>
<td>78</td>
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<tr>
<td></td>
<td>Medium</td>
<td>178</td>
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<tr>
<td></td>
<td>Dark</td>
<td>19</td>
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<tr>
<td>Gray-Brown</td>
<td>Light</td>
<td>4</td>
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<tr>
<td></td>
<td>Medium</td>
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<tr>
<td>Red-Brown</td>
<td>Light</td>
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<td></td>
<td>Medium</td>
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<tr>
<td></td>
<td>Dark</td>
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<tr>
<td>Red</td>
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<td></td>
<td>Dark</td>
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</tr>
</tbody>
</table>
(a) Archetype for colorless hair from this study sample. This hair was acquired from K2 B249.

(b) Archetype for light blond hair from this study sample. This hair was acquired from K2 B261.

(c) Archetype for blond hair from this study sample. This hair was acquired from K2 B44.

(d) Archetype for light golden brown hair from this study sample. This hair was acquired from K2 B281.

(e) Archetype for dark golden brown hair from this study sample. This hair was acquired from K2 B281.

(f) Archetype for light brown hair from this study sample. This hair was acquired from K2 B265.

(g) Archetype for dark brown hair from this study sample. This hair was acquired from K2 B81.

(h) Archetype for light gray-brown hair from this study sample. This hair was acquired from K2 B261.

Figure 3.7: Hair comparison archetype reference chart for Kellis 2 sample population.
Pigment Characteristics

Human beings are unique among primate species in that they are the only ones to grow long, thick, and highly-pigmented scalp hair. Some propose that this may have been an adaptation in response to a diet heavy in marine resources. Since these resources regularly contain a large number of harmful toxins “the selective and avid binding of toxins and metals to melanin within a rapidly growing and highly melanized tissue [such as hair] would have been an important adaptation” (Tobin 2008:234). Other possibilities for the excessive growth and pigmentation of human hair include sexual dimorphism and protection from environmental factors (Tobin 2008:234).

The latter possibility is sometimes used to explain the darker pigmentation generally attributed to individuals with African ancestry and those from tropical climates. Darker skin and hair are thought to be adaptations offering “protection against sunstroke afforded by deep skin and hair pigmentation as well as the contribution of melanin’s very efficient and fast exchange of ion transport and efflux for adequate salt balance” (Tobin 2008:234). According to Tobin (2008) a full 90% of the human population has “'environmentally friendly' brown-black hair and tanning skin”, while the remaining 5-10% has blonde or red hair and lighter skin pigmentation (234). In addition to melanin, when discussing pigment in human hair several other variables also need to be considered. These include pigment density, pigment distribution, pigment aggregate shape, and pigment aggregate size. These are discussed in detail in the following sections.
Pigment Density

Pigment density is defined by the *Atlas of Human Hair* (1999) as “the abundance of pigment granules observed using light microscopy” (28). Archetypes for absent, light, medium, and heavy pigment density acquired from the study sample can be seen in Figure 3.8 below.

(a) Archetype for absent pigment density. This hair was acquired from K2 B281.

(b) Archetype for light pigment density. This hair was acquired from K2 B281.

(c) Archetype for medium pigment density. This hair was acquired from K2 B131.

(d) Archetype for heavy pigment density. This is a beard hair acquired from K2 B81.

Figure 3.8: Pigment Density archetype reference chart for Kellis 2 sample population.
Pigment Distribution

Pigment distribution refers to how and where pigment is concentrated along the hair shaft. Pigment can be distributed uniformly, to one side of the hair shaft, peripherally, and randomly. In cases where the hair is colorless or pigment is unnoticeable, pigment distribution is marked as absent. Archetypes for pigment distribution can be seen in Figure 3.9 below.

(a) Archetype for absent pigment distribution. This hair was acquired from K2 B309.

(b) Archetype for uniform pigment distribution. This hair was acquired from K2 B131.

(c) Archetype for one-sided pigment distribution. This hair was acquired from K2 B135.

(d) Archetype for peripheral pigment distribution. This hair was acquired from K2 B261.

Figure 3.9: Pigment distribution archetype reference chart for Kellis 2 sample population.
Pigment Aggregate Shape

According to the *Atlas of Human Hair* (1999) “pigment aggregate shape refers to the appearance of the pigment granules when they are concentrated in a mass that has a recognizable form” (30). In other words, when individual melanin pigment granules coalesce they tend to form specific shapes including streaks and clumps or a mixture of both. Figure 3.10 represents archetypes for pigment aggregate shape.

(a) Archetype for clump pigment aggregate shape. This hair demonstrates large aggregate clumps. It was acquired from K2 B281.

(b) Archetype for streak pigment aggregate shape. This hair demonstrates small aggregate streaks. It was acquired from K2 265.

(c) Archetype for other (mixed) pigment aggregate shape. It was acquired from K2 B281.

Figure 3.10: Pigment aggregate shape archetype reference chart for Kellis 2 sample population.
Pigment Aggregate Size

Pigment aggregate size, as its name suggests, concerns the size of the pigment aggregates themselves. Figure 3.11 demonstrates small, medium, and large pigment aggregate size archetypes found within the study sample. In cases where pigment aggregates were not observed, pigment aggregate size was marked as absent.

(a) Archetype for small pigment aggregate size. This hair demonstrates small aggregate streaks. It was acquired from K2 B309.

(b) Archetype for medium pigment aggregate size. This hair demonstrates medium aggregate streaks. It was acquired from K2 B131.

(c) Archetype for large pigment aggregate size. This hair demonstrates large aggregate clumps. It was acquired from K2 B261.

Figure 3.11: Pigment aggregate size archetype reference chart for Kellis 2 sample population.
Medulla Characteristics

Put into the very simplest of terms, hair can be described as being shaped like a tube or a straw. Placed into the realm of biological anthropology, the structure of human hair is comparable to human bone. Like bone there is a hard-outer shell, in this case it is provided by the keratinized outer root sheath which is then followed by the inner root sheath. Finally, the cuticle and cortex surround an air or fluid filled space in the middle of the hair shaft, and like the marrow filled hollow of human bone this space is called the medulla.

In thicker animal hair, like that seen in horse manes, cat whiskers, or porcupine quills, the medulla can take up a large portion of the hair structure itself. However, in thinner hair, as is found in most humans, “the medulla—if present—generally comprises only a small percentage of [the fiber] mass” (Robbins 2002:50). The medulla in human hair can take many forms. In some cases, the medulla is absent due to how thin the human hair shaft can be, but where medullae are present they can be continuous, interrupted, fragmented, streaked, and on rare occasions a double medulla may even be present (Harding and Rogers 1999:17; Robertson 1999:80; Robbins 2002:50). In cases where the medulla has been interrupted or fragmented, “the medulla may be broken transversely at irregular intervals by cortical material (interrupted medulla) or it may only be present irregularly in very small amounts in the cortex (fragmented medulla)” (Harding and Rogers 1999:17). Additionally, when observed microscopically the medulla can appear opaque, as happens when it is filled with air, or translucent if it is filled with fluid. Some hair will even show a combination of both opaque and translucent medulla.
The *Atlas of Human Hair* (1999) defines a total of seven medulla continuity archetypes and four medulla opacity archetypes. However, given the lengths of the hair samples acquired for this particular study, some of the medulla continuity archetypes do not apply. Since the hair samples utilized in this study were cut to approximately two inches long, so that they could better fit onto the slides, having more than one medulla continuity type within a single hair was generally not observable. Therefore, for purposes of this paper the archetypes for medulla continuity were reduced to four archetypes, which can be seen in Figure 3.12. These include absent, fragmented, interrupted, and continuous. The archetypes provided by the *Atlas of Human Hair* (1999) for medulla opacity, however, were all observed within the sample. Therefore, excluding absent, the archetypes for medulla opacity within the study sample can be seen in Figures 3.13. These archetypes include opaque, translucent, and opaque/translucent.
As mentioned above, since the hair samples were cut to approximately two inches the archetypes for medulla continuity were reduced from seven types to four (Figure 3.12).

(a) Archetype for absent medulla. This hair was acquired from K2 B131.

(b) Archetype for fragmented medulla continuity. This hair was acquired from K2 B204.

(c) Archetype for interrupted medulla continuity. This hair was acquired from K2 B249.

(d) Archetype for continuous medulla continuity. This hair was acquired from K2 B293.

Figure 3.12: Medulla continuity archetype reference chart for Kellis 2 sample population.
Medulla Opacity

This study utilized polarized light microscopy to observe individual hair characteristics; this method is discussed in detail within the methodology section. Both opaque and translucent medulla were observed in the study sample, and the archetypes for translucent, opaque, and opaque/translucent medulla opacity can be seen in Figure 3.13.

(a) Archetype for transparent medulla opacity. This hair was acquired from K2 B44.

(b) Archetype for opaque medulla opacity. This hair was acquired from K2 B309.

(c) Archetype for other medulla opacity. The medulla for this hair is both opaque and transparent. This hair was acquired from K2 B293.

Figure 3.13: Medulla opacity archetype reference chart for Kellis 2 sample population.
**Cuticle Characteristics**

The cuticle is the hard, outer layer of the hair shaft. It is made up of scales, which interlock and overlap both longitudinally and laterally. The cuticle’s main purpose is to effectively hold the cortex together, while also protecting the cortex from chemical and environmental damage (Harding and Rogers 1999: 22; Robbins 2002:26). In mature human hairs, the cuticle is approximately six scales thick and the scale pattern, described using Wildman’s nomenclature, is ‘close wave’; although, it should be noted that scale patterns can vary slightly between individuals (Harding and Roger 1999:22; Robertson 1999:115; Morioka 2005:58). Scale pattern is especially important in the field of forensic hair examination since it has been shown to differ between species. This being the case, cuticle scale pattern can effectively distinguish human from non-human hairs.

However, since it has been demonstrated by previous researchers that cuticle scale pattern is very similar among humans, and since all of the hairs in this study sample were known to be of human origin, cuticle scale pattern was not included among the cuticle characteristics examined in the study sample. Instead, the research in this thesis is focused on cuticle characteristics that varied from one hair to the next. These characteristics include cuticle thickness (Figure 3.14), inner cuticle margin (Figure 3.15), and cuticle scale profile (Figure 3.16).
Cuticle Thickness

Cuticle thickness refers to how thick the cuticle appears, that is the thickness between the inner cuticle margin and the outermost edge of the cuticle. This can be subjective unless an accurate measure can be taken of the cuticle itself. However, there is a notable difference between thick and thin cuticle, which is why they are broadly defined as such for purposes of this thesis. Additionally, varied cuticle thickness was observed in the study sample. This occurs when the thickness of the cuticle does not remain constant along the hair shaft.

(a) Archetype for unknown cuticle thickness. This hair was acquired from K2 B281.

(b) Archetype for thin cuticle thickness. This hair was acquired from K2 B274.

(c) Archetype for thick cuticle thickness (space between black markers). This hair was acquired from K2 B22.

(d) Archetype for varied cuticle thickness. This hair was acquired from K2 B131.

Figure 3.14: Cuticle thickness archetype reference chart for Kellis 2 sample population.
Inner Cuticle Margin

The inner cuticle margin is usually distinguishable when the hair is heavily or moderately pigmented. In cases where the hair is lightly pigmented or colorless, the inner cuticle margin is generally indistinct. This is not always the case, but the figures below are a good example of this generality. A clear separation can be seen between the inner cuticle margin and the rest of the hair shaft for the more heavily pigmented hair seen in (a) of Figure 3.15, while the inner cuticle margin for the lightly pigmented hair in (b) cannot be distinguished from the rest of the hair shaft.

Inner Cuticle Margin Archetype Chart

(a) Archetype for distinct inner cuticle margin. This hair was acquired from K2 B135.  
(b) Archetype for indistinct inner cuticle margin. This hair was acquired from K2 B261.

Figure 3.15: Inner cuticle margin archetype reference chart for Kellis 2 sample population.
Cuticle Scale Profile

Cuticle scale profile refers to the appearance of the cuticle on the outside of the hair shaft. The most prolific archetype for cuticle scale profile seen in this study sample was smooth, while the least observed archetype was looped. Figure 3.16 provides archetypes for smooth, rough, and serrated cuticle scale profiles. In cases where the cuticle was severely damaged, the cuticle scale profile was listed as unknown.

(a) Archetype for smooth cuticle scale profile. This hair was acquired from K2 B81.

(b) Archetype for ragged cuticle scale profile. This hair was acquired from K2 B131.

(c) Archetype for serrated cuticle scale profile (black line shows outermost cuticle edge). This hair was acquired from K2 B.

Figure 3.16: Cuticle scale profile archetype reference chart for Kellis 2 sample population.
Ovoid Bodies

According to Robertson (1999) “ovoid bodies are well-defined, highly dense clumps of undispersed pigment” (113). They can vary in size and distribution. Some hairs can have many ovoid bodies, while others may have very few to none. Within the study sample the majority of the hairs examined did not show the presence of ovoid bodies. Some samples, however, did show many ovoid bodies and some showed few ovoid bodies. The archetypes for ovoid bodies are shown below in Figures 3.17.

(a) Archetype for absent ovoid bodies. This hair was acquired from K2 B.

(b) Archetype for few ovoid bodies. This hair was acquired from K2 B.

(c) Archetype for many ovoid bodies. This hair was acquired from K2 B281.

Figure 3.17: Ovoid bodies archetype reference chart for Kellis 2 sample population.
Sample Preparation

There were two methods of preparation, dry and wet, utilized in this study. For the dry method, hairs were extracted directly from the scalp sections of 51 individuals and placed in labeled bags. Conversely, for the wet method 1cm² scalp sections, acquired from ten of the original 51 individuals, were soaked for 24 to 48 hours in dimethyl sulfoxide (DMSO) to rehydrate the scalp following the method outlined in Williams (2008). Hairs were then extracted individually from the scalp and left to dry overnight. These two methods were used to help determine if sample preparation also affects PHRM and to help initiate standardization procedures for analysts examining PHRM. Results for this portion of the study can be seen in Chapter 2 of this thesis.

After all of the samples were procured, four hairs were mounted onto each labeled glass slide using Permount®; for a total of 5-10 slides per individual depending on the preparation method used. Permount® was chosen as a mounting medium because it has a refractive index (RI) of 1.525 at 25°C, which is very similar to the RI of human hair at 1.55. This makes it an optimal mounting medium for this project (Linch and Prahlow 2001; Marschner et al. 2003; Committee on identifying the needs of the forensic sciences community 2009:156). After the hairs were mounted and the slides were fully dried, each slide was observed using an Amscope 40x-1500x Infinity Polarizing Microscope at 40-60x magnification.

Results

Using Chi-square analyses, each of the intrinsic factors listed above were analyzed with SPSS® statistics software to determine their correlation to PHRM. Unsurprisingly, the correlation between PHRM and growth stages, as demonstrated through previous
research (Petraco et al. 1988; Linch and Prahlow 2001; Jeffery et al. 2007; Koch et al. 2013), proved to be significant at \( p=.000 \). Again, this was not surprising; several studies have reported a link between putrid roots and growth stages. There were several other intrinsic factors, however, that demonstrated statistical significance in correlation with PHRM including age, cuticle thickness, cuticle scale profile, and hair color. The remaining variables did not show a statistically significant correlation (\( p\geq 0.05 \)). Inner cuticle margin, medulla opacity, ovoid bodies, medulla continuity, pigment aggregate shape, pigment distribution, biological sex, and pigment aggregate size appear to be unrelated to the incidence of putrid roots, though further studies are needed to rule out a relationship entirely. Table 3.2 provides the statistical analysis for these variables in relation to PHRM.

Table 3.2: Pearson Chi-Square results for intrinsic characteristics not demonstrating a significant correlation with changes in postmortem hair root morphology.

<table>
<thead>
<tr>
<th></th>
<th>Asymp. Sig. (2-sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inner Cuticle Margin</td>
<td>.070</td>
</tr>
<tr>
<td>Medulla Opacity</td>
<td>.109</td>
</tr>
<tr>
<td>Ovoid Bodies</td>
<td>.125</td>
</tr>
<tr>
<td>Medulla Continuity</td>
<td>.181</td>
</tr>
<tr>
<td>Pigment Aggregate Shape</td>
<td>.417</td>
</tr>
<tr>
<td>Pigment Distribution</td>
<td>.644</td>
</tr>
<tr>
<td>Biological Sex</td>
<td>.807</td>
</tr>
<tr>
<td>Pigment Aggregate Size</td>
<td>.864</td>
</tr>
</tbody>
</table>
Growth stages, as mentioned above, demonstrated a correlation with PHRM with p=.000 (Table 3.3). The greatest proportion of postmortem changes were seen in hairs in anagen stage, with very few changes occurring within the telogen stage. Nearly 43% (118/277) of anagen roots showed postmortem changes in hair root morphology, with the majority of these showing evidence of PMRB at over 70% (86/118) of the total changes observed. Thirty hairs also had BLE and four were a mix of BLE and PMRB. The remaining two hairs demonstrated changes that are consistent with HKP, a postmortem change usually associated with wet decomposition and not something customarily attributed to dry desert environments. Figure 3.18 is a bar graph detailing the distribution of postmortem changes in association with hair growth stages.

Table 3.3: Pearson Chi-Square results for correlation between growth stages and root decomposition.

<table>
<thead>
<tr>
<th>Value</th>
<th>Asymp. Sig. (2-sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pearson Chi-Square</td>
<td>274.689a</td>
</tr>
<tr>
<td>N of Valid Cases</td>
<td>713</td>
</tr>
</tbody>
</table>

a. 0 cells (0.0%) have expected count less than 5. The minimum expected count is 8.46.
Figure 3.18: Bar graph showing the proportion of postmortem changes seen within the sample population and how they are correlated with growth stages.
Age also demonstrated statistical significance at p=.000 (Table 3.4), with those in the youngest age group (16-21) showing the greatest correlation; 56% (19/34) of individuals in this age range exhibited changes in PHRM. The lowest incidence of change is in individuals between the ages of 36-50 (20%, 30/149). And, while it should be noted that a large portion of the study population lies within this age range, the greatest proportion of individuals were aged between 22-35 and they demonstrated the second largest correlation at 31% (91/290). It was also determined that individuals in the older age ranges (51-60 and 60+) showed evidence of putrid roots in only 25% (61/240) of the population, suggesting the possibility that the occurrence of putrid roots decreases as people age. It would be interesting to see if the same would hold true with a larger sample population. Figure 3.19 shows a bar graph of postmortem changes in hair root morphology and their relation to age.

Table 3.4: Pearson Chi-Square results for correlation between age and root decomposition.

<table>
<thead>
<tr>
<th></th>
<th>Value</th>
<th>Asymp. Sig. (2-sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pearson Chi-Square</td>
<td>20.061a</td>
<td>.000</td>
</tr>
<tr>
<td>N of Valid Cases</td>
<td>713</td>
<td></td>
</tr>
</tbody>
</table>

a. 0 cells (0.0%) have expected count less than 5. The minimum expected count is 9.58.
Figure 3.19: Bar graph showing the proportion of postmortem changes within the sample population and how it correlates with age.
The next variable that appears to be associated with postmortem changes in hair root morphology is cuticle thickness. At p=.002 (Table 3.5) cuticle thickness seems to be nearly as significant as growth stages and age to the formation of putrid roots within this study population. Hairs with thick cuticles comprise nearly 30% (206/713) of the sample, but are ~18% more likely to exhibit postmortem changes. Additionally, 74% (52/70) of the samples with thick cuticles that show evidence of postmortem changes has PMRB. This is in opposition to the pattern exhibited by thin and varied cuticles. In thin and varied cuticle thickness, the majority of postmortem changes observed included brush-like ends (62%, 13/21, and 71%, 25/35, respectively). Unfortunately, one-third (237/713) of the sample included hairs with unknown cuticle thickness. Generally, these were hairs with light pigment density or those that were entirely colorless (see Figure 3.20). Despite this, it is interesting to note that hair with varied cuticle thickness lies somewhere in the middle when it comes to showing evidence of postmortem changes (25%, 35/142). This suggests that there is likely to be a correlation between cuticle thickness and changes in PHRM. Thin cuticles showed evidence of postmortem changes in only 16% (21/128) of the hairs studied. Figure 3.21 is a bar graph detailing the relationship between cuticle thickness and postmortem changes in the hair root.
Table 3.5: Pearson Chi-Square results for correlation between cuticle thickness and root decomposition.

<table>
<thead>
<tr>
<th></th>
<th>Value</th>
<th>Asymp. Sig. (2-sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pearson Chi-Square</td>
<td>14.470&lt;sup&gt;a&lt;/sup&gt;</td>
<td>.002</td>
</tr>
<tr>
<td>N of Valid Cases</td>
<td>713</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> 0 cells (0.0%) have expected count less than 5. The minimum expected count is 36.08.

Figure 3.20: A colorless hair from K2 B293 demonstrating an unknown cuticle thickness.
Figure 3.21: Bar graph showing the proportion of postmortem changes documented within the sample population and how they are correlated with cuticle thickness.
A link between changes in PHRM and pigment density was also statistically significant at $p=.006$ (Table 3.6). PMRB had the greatest representation with 46% (55/119) of these cases occurring in hairs with a medium pigment density. Additionally, the majority of hair exhibiting postmortem changes occurred in this category (35%, 81/230). Unfortunately, heavily pigmented hair was not well represented in the sample population; only 6% (41/713) of hairs were heavily pigmented, while the great majority of hairs were lightly pigmented (37%, 265/713), or medium pigmented (32%, 230/713). This could be partially due to old age- 55% (389/713) of the study population was aged 36 or over- but also because the sample was comprised of specimens from a single locale suggesting probable physical and genetic similarities between individuals. As suggested earlier, a greater diversity within the study sample would help to elucidate these results. The line graph in Figure 3.22 provides a visual representation of the relationship between pigment density and PHRM. It is clearly demonstrated that PMRB occurs the most in hairs with medium pigment density.

Table 3.6: Pearson Chi-Square results for correlation between pigment density and root decomposition.

<table>
<thead>
<tr>
<th></th>
<th>Value</th>
<th>Asymp. Sig. (2-sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pearson Chi-Square</td>
<td>12.548a</td>
<td>.006</td>
</tr>
<tr>
<td>N of Valid Cases</td>
<td>713</td>
<td></td>
</tr>
</tbody>
</table>

a. 0 cells (0.0%) have expected count less than 5. The minimum expected count is 11.56.
Figure 3.22: Bar graph showing the proportion of postmortem changes documented within the sample population and how they are correlated with pigment density.
Similarly, when regarding cuticle scale profile (CSP), a lack of diversity could be to blame for the overwhelmingly smooth cuticles seen in this sample. Out of 694 individual hairs where CSP could be discerned, a full 426 (59%) of them had smooth CSP, while ragged accounted for ~10% (69/694), serrated 3% (22/694), and looped <.01% (4/694). While those hairs with smooth CSP did demonstrate a closer association with changes in PHRM, they were 16% more likely to show postmortem changes than those hairs with ragged CSP-this could be due to the lack of diversity within the study sample. Therefore, the statistical analysis regarding CSP is likely skewed. Further research should include broadening the scope of the study population.

Table 3.7: Pearson Chi-Square results for correlation between cuticle scale profile and root decomposition.

<table>
<thead>
<tr>
<th></th>
<th>Value</th>
<th>Asymp. Sig. (2-sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pearson Chi-Square</td>
<td>18.116(^{a})</td>
<td>.003</td>
</tr>
<tr>
<td>N of Valid Cases</td>
<td>713</td>
<td></td>
</tr>
</tbody>
</table>

\(^{a}\) 4 cells (33.3%) have expected count less than 5. The minimum expected count is .28.
Figure 3.23: Bar graph showing the proportion of postmortem changes documented within the sample population and how they are correlated with cuticle scale profile.
The last intrinsic characteristic that showed a statistically significant correlation with putrid roots is hair color. Fortunately, unlike CSP, hair color was widely represented. The most abundant color was medium brown, which accounted for 25% (178/713) hairs, but dark golden brown (20%, 145/713) and colorless hair (18%, 130/713) were also present in large quantities. The least represented colors were medium gray-brown (1), medium red (1), dark red (3), and light gray-brown (4). Considering that red hair occurs in only about 2% of the world population, it is not surprising that few exist in the study sample. However, because there were a few hair colors represented by small numbers, analysis was run using SPSS® statistics software on all hair colors as well as on hair colors with 50 or more representations in the sample; this latter analysis was comprised of 597 hairs. Hair colors with a sample of 50 or more samples include colorless, dark golden brown, light blonde, light brown, and medium brown.

After excluding colors that were not well represented, the analysis gave a p-value of .006 (Table 3.9), showing a greater statistical significance than the p-value of .013 when all hair colors were included into the analysis (Table 3.8). Each hair color demonstrated a similar propensity for changes in PHRM (~30%-35%) except for light brown hair. The incidence of putrid roots in light brown hair was reduced to a low 11% (9/78), a significant reduction from the amount seen in other hair colors. Unfortunately, it is not understood why light brown hair would show fewer changes, but perhaps this goes hand in hand with cuticle thickness. More than half of the light brown hairs had thin or varied cuticles, which, as mentioned earlier, accounts for fewer cases of putrid roots than those hairs with thick cuticles. Contrary to this, dark golden brown hairs, which showed the greatest proclivity for putrid roots, had a larger percentage of thick cuticles.
Table 3.8: Pearson Chi-Square results for correlation between hair color and root decomposition.

<table>
<thead>
<tr>
<th></th>
<th>Value</th>
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<tr>
<td>Pearson Chi-Square</td>
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<td>.013</td>
</tr>
<tr>
<td>N of Valid Cases</td>
<td>713</td>
<td></td>
</tr>
</tbody>
</table>

a. 12 cells (37.5%) have expected count less than 5. The minimum expected count is .28.

Table 3.9: Pearson Chi-Square results for correlation between hair colors represented by 50+ hair samples and root decomposition.

<table>
<thead>
<tr>
<th></th>
<th>Value</th>
<th>Asymp. Sig. (2-sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pearson Chi-Square</td>
<td>14.636</td>
<td>.006</td>
</tr>
<tr>
<td>N of Valid Cases</td>
<td>597</td>
<td></td>
</tr>
</tbody>
</table>

a. 0 cells (0.0%) have expected count less than 5. The minimum expected count is 18.79.

Discussion

As mentioned earlier, it is not surprising to find an association between hair root growth stages and PHRM. This characteristic has been studied by several researchers (Petraco et al. 1988, Linch and Prahlow 2001, Koch et al. 2013) and it has already been established that anagen and catagen stage hairs are more susceptible to postmortem changes than telogen hairs. Again, the full keratinization of telogen hair roots likely causes its resistance to postmortem root degradation.
Age related changes to hair keratin also explains why age affects PHRM. Younger individuals have less cysteine disulphide in their hair, which means “children’s hair has lower hardness… than adult hair” (Tang et al. 2015: 160-161). Cysteine disulphide “forms cross-linkages in cuticle cells and contribute to the physical and mechanical properties of hair” (Tang et al. 2015: 160). Therefore, younger hair, being deficient in cysteine disulphide, would seem to be more vulnerable to postmortem degradation. This assumption is bolstered by the analysis which shows that individuals in the age range of 16-21 are more prone to decomposition changes in their hair than older individuals.

The greater proportion of PMRB seen in thick cuticles can probably be explained by looking at the greater percentage of BLE seen in thin cuticles. A thinner cuticle could cause the hair to be more easily broken when forcibly removed from the scalp. Since PMRB causes the root end to be more fragile, a thinner cuticle is more likely to break in this area thereby resulting in a greater proportion of BLE. Thicker cuticles would be less likely to break in the area of PMRB, meaning thick cuticles would be more inclined to show evidence of PMRB than BLE.

The final three intrinsic factors and their affinity for PHRM can be explained by limited sample size and limited sample diversity. Pigment density was similar throughout the sample, with most samples showing a light or medium pigment density; this can partially be due to the overall older age profile of the individuals included in the study population. Unfortunately, heavy pigment density was not adequately represented. A larger, more diverse sample population would help determine whether pigment density affects PHRM. Cuticle scale profile and hair color would similarly benefit from a more diverse sample population. 426/694 hairs, where CSP could be discerned, has smooth
CSP; that is equal to 59% of the total sample. It is fair to assume that this probably skewed the results. The similarity of hair colors, likely due to similar ancestry, probably biased the results in an identical manner.

**Benefits and Limitations of Research**

Now, more than ever, the admissibility of certain evidence is under scrutiny. Among these is the utility and reliability of forensic analysis involving human hair. This comes in the wake of an article by *The Washington Post* in which it was stated that, “the Justice Department and FBI have formally acknowledged that nearly every examiner in an elite FBI forensic unit gave flawed testimony in almost all trials in which they offered evidence against criminal defendants over more than a two-decade period before 2000” (Hsu 2015). This included “overstat[ing] forensic matches in ways that favored prosecutors”; in 32 of those cases the defendant was sentenced to death (Hsu 2015:1). While it is unlikely that this evidence alone resulted in a conviction, it does raise questions about the authenticity of forensic analyses overall.

Fortunately, most types of admissible forensic evidence have proven their efficacy over the course of history. The reliability of fingerprints and DNA are rarely questioned, and blood spatter and ballistics have both shown to be dependable in criminal investigations. With the introduction of television series such as *CSI, Bones,* and *NCIS,* jurors are beginning to think of themselves as armchair detectives, and therefore are requiring more evidence as a result. Furthermore, those who watch these crime procedurals are led to believe that forensics is an exact science that can be accepted without question, that evidence can speak for itself, and that the methods of crime scene investigators are infallible (Kruse 2010:82-88). This means that the science behind
admitted evidence needs to be sound. Firmly put, if PHRM is better understood and if its causes can be determined, the potential impact for medico-legal cases could be enormous.

However, there are several limitations to studies such as this. The main limitation is sample size. In order to acquire the large and diverse sample necessary to understand the causes of putrid roots, cadavers from several different ancestries, both sexes, and various ages need to be represented in the sample population. Additionally, bodies should be placed in diverse, but controlled, environments and studied at various stages of decomposition. Climate, predatory activity, insect populations, fungal growth, and other extrinsic factors also need to be considered in determining the variables included within the sample set. Unfortunately, the likelihood of a research sample with these criteria is improbable. Therefore, it is necessary to undertake a great amount of research in order to provide analysts with the best data regarding PHRM.

Conclusion

There are several hurdles to overcome before PHRM can be accepted as reliable evidence. In addition to the issues stated above, there is also the problem of linking a questioned hair to a known source. Without a way to identify the hair as belonging to a specific deceased individual, PMRB, BLE, and HKP become moot points. Does the appearance of changes in hair root morphology suggest decomposition? More than likely. But without an understanding of the causes behind these changes, PMRB, BLE and HKP bring minimal evidentiary value to criminal cases. Finding significant correlations between PHRM and intrinsic variables, therefore, provides a better understanding of the mechanisms involved in postmortem hair root degradation.
This study shows that there are more than extrinsic variables acting upon PHRM. Much like decomposition of the body itself, decomposition in hair is influenced by intrinsic factors. Age showed a greater association with hair root degradation, as did cuticle thickness, pigment density, cuticle scale profile, and hair color. Both age and cuticle thickness and their associations with PHRM make sense. Younger individuals have softer and less structurally sound hair than older individuals due to the lack of cysteine disulphide in their hair. This makes their hair and hair roots more susceptible to postmortem changes. Additionally, in regards to cuticle thickness, it makes sense that thinner cuticles are more likely to lead to BLE. A thinner cuticle means less support for the hair shaft and a greater likelihood of breakage when the hair is forcibly removed from the scalp. This statement is supported by the analysis which shows that thinner roots showed more evidence of BLE, while thicker roots showed more evidence of PMRB. Unfortunately, while growth stages, age, and cuticle thickness are well represented within the sample, pigment density, cuticle scale profile, and hair color were skewed. These will require further study in order to discover their association with PHRM.

The ultimate aim of this research is that future forensic analysts will use this study as a stepping-stone, and broaden the scope of studies on PHRM. While hair analysis is still an imperfect science, and it has proven to be more so since fabrication of such evidence has come to light, hopefully more research will help to bolster its credibility. Further research should look at those intrinsic factors that did show an association, however weak, and attempt to discern whether there is in fact a connection. Additionally, other researchers should look at those factors without strong connections, such as
medulla continuity and medulla opacity, to determine why these do not have an impact on PHRM.
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Molto, J.E.

Molto, J.E.

Morioka, K.

Ogle, Jr Robert, Michelle J. Fox.

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Robbins, Clarence R.

Robertson, James.


Williams, Lana J.  

Wollina Uwe.  
CHAPTER 4: CONCLUSION

While other animals sport feathers and scales, mammals alone grow hair. It enables them to keep warm, and for some serves as a useful scare tactic. For humans, however, the necessity and overall usefulness of hair is still questioned. Unlike our mammalian counterparts, for most of us hair is used more as an accessory. We style it, color it, cut it, and in some cases, we shave it all off.

Fortunately, although we may not know the bodily or evolutionary function for the retention of human hair, studies in hair analysis have taught us that human hair itself is unique. It has a specific pattern, like a fingerprint, that sets it apart from hair in other mammalian species. Because all mammals have hair that differs structurally from that of other mammals, analysts are able to differentiate between species when examining a single hair. This, along with other applications, have become useful in the field of forensics, where the questioning of known and unknown hairs, DNA analysis, heavy metal analysis, and toxicological analysis can mean the difference between catching the guilty party or even ensuring an innocent person is not convicted for a crime they did not commit. One such forensic application has been the subject of this thesis. First, two preparation methods were assessed to determine their efficacy when investigating mummified remains. Then the evidentiary significance and reliability of postmortem human hair degradation in regards to medico-legal investigations were evaluated by determining which intrinsic variables, if any, may have an effect on the occurrence of degradation.
Hair samples acquired from fifty individuals from the Kellis 2 cemetery in the Dakhleh Oasis, Egypt, were prepared using two preparation methods. These included both a wet and dry method, and the resulting samples were evaluated to determine the efficacy of each method. It was determined that the wet method, which involved rehydrating the tissue using DMSO, provided the most definitive and advantageous results. The hair samples acquired from the rehydrated scalp sections were more easily obtained, and therefore resulted in less breakage and a better representation of anagen and catagen hair roots. As postmortem hair degradation is generally relegated to these earlier stages of hair growth, it is crucial that a large number of these are collected to provide a more significant study sample for investigation.

Additionally, fewer roots were lost during the acquisition of hairs from the rehydrated scalp sections. The breakage of the hair root from the shaft occurred nearly half as often in the wet samples as it did in the dry samples (20% and 36% respectively). Lastly, the occurrence of postmortem hair degradation and types of degradation documented was significantly different between the two preparation methods. In the dry samples, there were equal incidences of both BLE and PMRB, while in the wet sample there was a predominance of PMRB (25%) within the sample and BLEs made up only 3%. There are two possible explanations for this disparity. The least likely explanation is that there simply were more BLEs in the dry sample, but more than likely the cause of the imbalance resulted from the forceful removal of the hair root from a dry scalp. This could cause the root to break off, thereby producing a BLE on the proximal end of the shaft.

While there are some studies (e.g., Petraco et al. 1988; Tafaro 2000; Linch and Prahlow 2001; Domzalski 2004; Jeffery et al. 2007; Koch et al. 2013) that examine how
extrinsic variables such as time since death, climate, method of decomposition, stage of decomposition, and burial environment affect postmortem hair root morphology, none have seriously considered intrinsic factors. It is already widely believed that postmortem hair degradation proceeds in a predictable manner (e.g., Petraco et al. 1988; Tafaro 2000, Linch and Prahlow 2001; Domzalski 2004; Jeffery et al. 2007; Koch et al. 2013), where PMRB occurs in the earlier stages of decomposition, and BLEs and HKPs are thought to be related to later stages of dry and wet decomposition respectively. This study calls this into question. The individuals in this sample have been buried since AD50-AD450 in a hyper arid environment, and PMRB was present in the sample as well as HKP, which are thought to be related to moist decomposition. While the HKP may be able to be explained by the fact that different portions of the body may be subjected to different types of decomposition within a single burial environment, this does not explain why PMRB was found on hairs from individuals who have been deceased for nearly 1600-2000 years. Therefore, if time since death, climate, method of decomposition, stage of decomposition, and burial environment do not fully explain why postmortem hair degradation occurs, other factors need to be taken into account.

The second portion of this thesis concerns the effect that intrinsic variables might have on the incidence of postmortem hair degradation. A total of 713 hairs were examined for color, pigment density, pigment distribution, pigment aggregate shape, pigment aggregate size, medulla continuity, medulla opacity, cuticle thickness, inner cuticle margin, cuticle scale profile, and ovoid bodies. Additionally, age and biological sex were recorded for each individual and included as part of the intrinsic variables.
While not all of these variables showed a significant correlation with postmortem modification of the hair root, several did. The growth stage of the hair appeared to have a significant impact on postmortem changes. Only roots in anagen and catagen growth stages showed these changes. Hair roots in the telogen growth stage did not demonstrate any of these modifications. This finding is in keeping with studies conducted by previous researchers within the field of hair analysis (e.g., Petraco et al. 1988; Tafaro 2000; Linch and Prahlow 2001; Domzalski 2004; Jeffery et al. 2007; Koch et al. 2013). In addition to growth stage, however, age, cuticle thickness, pigment density, cuticle scale profile, and hair color were shown to have a significant correlation with the appearance of postmortem degradation in the human hair root. This suggests that there are other factors affecting postmortem hair root degradation in addition to those that have been previously studied.

**Benefits and Limitations of Research**

The limitations to studies such as this are many and include: the need for a large sample placed under controlled circumstances and regularly monitored; the need for a large number of cadavers with diverse backgrounds and ancestry; and the need for samples to be observed in numerous environmental conditions. This study was very limited in scope since it was conducted on a single population of individuals who were buried in similar, if not the same, burial environment, and their bodies all underwent natural mummification. Additionally, the scale of this study was limited to 51 individuals and the decomposition processes were not regularly monitored. This means that the decomposition process may have been affected by factors which are not known. As such, this study can be considered to be a pilot project.
Despite this, there are several benefits to this study. For one, it shows that PMRB does not appear to be dependent on time since death. These individuals have been dead for over 1500 years, yet some hairs still show evidence of PMRB. It also provides a good representation of postmortem hair root changes in naturally mummified remains and remains buried in arid environments. Furthermore, the methodology section provides a good demonstration of preparation methods for hair samples from mummified remains. This will hopefully lead to the standardization of hair analysis preparation methods in future research.

**Future Considerations**

For postmortem hair degradation to be widely accepted as evidence in medico-legal cases, more rigorous experimentation and research is needed. This thesis covers merely a fraction of the scenarios that can affect human hair roots after death. Neither the scope nor the scale of this investigation is large enough to encompass the entirety of situations that affect human decomposition. In order to cover such a diverse set of circumstances, a large-scale investigation would need to be conducted taking into account climate and environmental variability, predation, burial environments, manner and cause of death, underlying medical conditions, intrinsic variables such as those listed above, and other factors that may affect postmortem hair root morphology.

Hair exhibiting PMRB is useless to investigators if it cannot be stated that it is undoubtedly attributable to the presence of decomposition. While Lowe (*State of Florida v. Casey Marie Anthony*, 48-2008-CF-015606-O 2011) did state that she had not come across any instances of PMRB where the individual was not deceased, questions still
arise about the veracity of this testimony simply because of the scarcity of published research.

Additionally, the question as to why PMRB occurs in some individuals while being absent in others needs to be resolved. This can only be accomplished with more research looking at extrinsic and intrinsic factors that affect PHRM. To lend credence to claims that PMRB and other postmortem morphological changes are the result of decomposition, and thereby are proof that a deceased individual was present, these studies need to be undertaken by experts within the fields of trace evidence, forensics, and hair analysis. Furthermore, sample sets need to become increasingly diverse, specimens need to be regularly monitored, and dispositional environments need to be strictly controlled. Future court cases will only benefit from this type of research.
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