Global downregulation of pigmentation-associated genes in human premature hair graying

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Abstract. Premature hair graying, or canities, is a complex multi-factorial process with negative effects on affected individuals. The aim of the present study was to investigate the possible underlying mechanisms of premature hair graying at the genetic level. A total of 5 unrelated Han Chinese individuals presenting with premature hair graying (25-40 years old, with >1% hair affected) were enrolled in the present study. RNA sequencing was performed to identify gene expression changes between the follicular cells of grey and black hair from the cohort. A total of 127 differentially expressed genes (DEGs) were identified. These DEGs were overrepresented in categories associated with the pigmentation pathway, with a decreased expression of key genes responsible for melanin synthesis. Of note, the decreased expression of certain transcription factors and the increased expression of certain precursor microRNAs observed may explain for the downregulation of certain other DEGs, which were identified as their targets via Starbase v2 and Integrated Motif Activity Response Analysis. The DEGs were also enriched in terms associated with the nervous system, indicating that neural disturbances may also have certain roles in premature hair graying. Of note, five of the downregulated DEGs were associated with aging according to the JenAge Aging Factor Database. To the best of our knowledge, the present study was the first genome-wide survey of the gene expression profile associated with premature hair graying. Dysfunction of the melanin biosynthesis pathway is probably the direct cause of hair graying and the present results provide valuable clues for further functional and mechanistic investigation.

Introduction

Hair graying may be classified as natural senile canities and premature graying. Natural senile canities usually has its onset after the age of 40 years and aggravates with the ongoing aging process (1). Unlike senile canities, premature graying occurs prior to the age of 25 years and is usually progressive and permanent (2,3). Although hair graying is considered to be genetically controlled and inheritable, the underlying mechanisms have remained largely elusive (4).

Hair color is determined by pigment granules in hair follicles, wherein melanin synthesis is particularly crucial (5). Mature melanocytes are densely distributed in hair bulbs to sustain active melanogenesis, which is strictly coupled to the anagen stage of the hair cycle (6,7). The biosynthesis of melanin and its subsequent transfer from melanocyte to hair bulb keratinocytes is a rather complex process (8). Previous molecular study has mainly focused on identifying genes that encode characteristic markers for pigmentation, including tyrosinase (TYR), OCA2 melanosomal transmembrane protein, TYR-related protein 1 (TYRPI) and solute carrier family 45 member 2 (SLC45A2) (9). Polymorphisms within these loci are associated with a normal variation in hair color traits (10). In addition, a genome-wide association study on the genetic basis of pigmentation in human subjects revealed that the single nucleotide polymorphism frequency distribution of these genes is linked to skin color and hair color (11,12). A broader study indicated that mutations in two pore segment channel 2, agouti signaling protein and melanocortin 1 receptor are associated with hair color and pigmentation (13).

Previous studies have focused on identifying genes that encode characteristic markers for premature graying of hair. A germline mutation in the syntaxin 17 gene of horses was recently identified to cause premature graying of hair (14) and telomerase reverse transcriptase mutation carriers may also present with premature hair graying (15). Reverse transcription polymerase chain reaction (RT-PCR) arrays on the gene expression profiles of the hair bulge and hair bulb revealed a significant downregulation of melanogenesis associated genes [tyrosinase (TYR), tyrosine related protein 1 (TYRPI),

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melanocyte inducing transcription factor, paired box gene 3 (PAX3) and proopiomelanocortin) in unpigmented hair bulbs and a downregulation of marker genes typical for melanocyte precursor cells [PAX3, SRY-Box 10 (SOX10) and dopachrome tautomerase] in unpigmented mid-segments compared with their pigmented analogues. Superoxide dismutase 3 transgenic mice exhibited premature aging, including hair graying, abnormal gait and a shortened life span (16).

Although previous studies provided candidate genes associated with hair graying, their results are limited due to the unavailability of related hereditary family members or an insufficient number of subjects to provide representative results (17). According to the central dogma of genetics, the genotype affects the phenotype through gene expression. Thus, it is straightforward to assume that a variance in gene expression between the grey and black hair follicles underlies the difference in hair color. However, to the best of our knowledge, no previous study has assessed premature hair graying from this genetic aspect. In the present study, an RNA sequencing (RNA-seq) analysis was performed to reveal gene expression changes between grey and black hair follicles from Han Chinese patients suffering from premature hair graying. It was intended to unravel the underlying mechanisms and potential candidate genes responsible for hair pigmentation loss.

Materials and methods

Subjects. A total of 5 unrelated Han Chinese donors who presented with premature graying since they were teenagers and had a clear family history of premature hair graying were enrolled in the present study. Their details are specified in (Table I). A total of 30 grey and 30 black hair follicles were randomly collected from each subject and stored in RNAlater® Stabilization Solution (Thermo Fisher Scientific, Inc., Waltham, MA, USA) at 4°C.

RNA extraction, library preparation and sequencing. Total RNA was extracted from each sample using the mirVana™ miRNA Detection Kit (Thermo Fisher Scientific, Inc.) in strict accordance with the manufacturer's protocol. The RNA-seq library was prepared using an Ion Total RNA-seq Kit v2 (Thermo Fisher Scientific, Inc.) following the manufacturer's protocol. RNA-seq libraries were sequenced on an Ion Proton™ System (Thermo Fisher Scientific, Inc.).

Bioinformatics analysis of RNA-seq data. Raw sequencing reads were aligned to the human genome (GRCh38/hg38) using STAR (Spliced Transcripts Alignment to a Reference software; http://code.google.com/p/rna-star/) (18). Ensembl gene annotation (http://www.ensembl.org/info/ontology/annotation.html; release 79) was used for evaluating gene expression using Cufflinks (19), and differentially expressed genes (DEGs) between grey and black hair follicles were determined by Cuffdiff, a subpackage of Cufflinks. The ratio of Fragments per kilobase per million of grey to black hair was calculated and differential expression genes were then studied by log2FC. The criteria used to define DEGs were as follows: i) Fragments per kilobase per million mapped reads >1 in at least one sample of a sample pair; ii) fold change of at least 2 and iii) P<0.05.

Functional classification and enrichment analysis of the DEGs were performed using the online platform of the Database for Annotation, Visualization and Integrated Discovery (20). Gene Ontology (GO) terms were used to classify the DEGs. Association between microRNAs (miRs) and their targets were predicted by Starbase v2 (21). Transcription factors (TFs) and their target genes were predicted through Integrated Motif Activity Response Analysis (22). The Search Tool for the Retrieval of Interacting Genes/Proteins (STRING) database (23) was used to construct protein-protein interaction (PPI) networks the DEGs are involved in.

RT-quantitative (q)PCR. Total RNA was extracted from each hair follicle sample. The RNA (1 µg) was reverse-transcribed using a ReverTra Ace qPCR RT Kit (Toyobo, Osaka, Japan). β-actin was used as reference control. SYBR Green Realtime PCR Master Mix (ROX; Toyobo) was applied for PCR amplifications on a Bio-Rad CFX96 system (Bio-Rad Laboratories, Inc., Hercules, CA, USA) according to the manufacturer's protocol. The thermocycling program was as follows: 95°C for 5 min, followed by 40 cycles of 95°C for 30 sec, 60°C for 40 sec and 72°C for 45 sec. The primers used are listed in Table II.

Results

Summary of RNA-seq data. To identify changes in the gene expression profile associated with premature hair greying, RNA-seq libraries derived from each grey or black hair follicle were sequenced using the Ion Proton™ System (Thermo Fisher Scientific, Inc.). The number of raw sequenced reads for each sample ranged from 15.9 to 27.9 million, and the mapping percentages were 72.1-80.0%, which demonstrated an overall good mappability.

Gene expression changes between grey and black hair follicles. In order to identify genes with a role in premature hair greying, the samples were classified into a grey hair group and a black hair group, and the gene expression changes were compared between them. With the criteria of a 2-fold change in expression and P<0.05, a total of 127 DEGs were identified, including 47 upregulated and 80 downregulated DEGs (Fig. 1A). In order to confirm these DEGs, the changes in their expression were also assessed for each subject individually. Most of these DEGs were changed in the same direction among the individual subjects (Fig. IB), demonstrating a consistency regarding the DEGs identified.

To determine the functional roles of these DEGs, the functional categories they were involved in were then examined. Of note, the DEGs were mainly enriched in functional categories closely linked to pigment synthesis or metabolic pathways (Fig. 1C). The DEGs were also known to be located in pigment granules and melanosomes (Fig. 1D), which provided an explanation for the downregulation of these genes in grey hair follicles that had lost their ability to produce melanin, which lead to hair graying. Of note, genes associated with the nervous system accounted for the largest number of DEGs, although the significance of their enrichment was not as high as that in the categories associated with pigmentation. This may imply a disturbance of the nervous system in grey hair follicles compared with that in black ones.
Genes associated with pigmentation and melanin synthesis are predominantly decreased in grey hair follicles. In order to obtain a comprehensive view of genetic changes associated with pigmentation and melanin synthesis, the DEGs identified were classified into pigmentation/melanin-associated GO terms (24). The results indicated that all of the pigmentation/melanin-associated DEGs were decreased in grey hair follicles (Fig. 2A). This suggested that pathways involved in melanin biosynthesis are downregulated in grey hair follicles. These DEGs comprised various genes that are well-known to be responsible for melanin biosynthesis, including TYR, melan-A (MLANA), premelanosome protein (PMEL), TYRP1, SLC45A2, KIT, G protein-coupled receptor 143 (GPR143) and OCA2.

Differentially expressed miRs and TFs regulating pigmentation/melanin-associated DEGs in grey hair follicles. Among the DEGs identified, 13 genes encoded miRs, all of which were significantly increased in grey hair follicles; they accounted for 28% (13/47) of all upregulated DEGs. It has been previously reported that miRs may be transcribed by RNA polymerase II and have polyA tails (25), and it was therefore likely that the miR-associated RNA-seq reads are from precursors of miRs. Since the majority of the DEGs (80/127; 63%) were decreased in grey hairs, the changes in the expression of DEGs targeted by these upregulated miR-encoding genes were examined (Fig. 2B). Of note, most of the miR-encoding DEGs (11 out of 13) had the other DEGs as their predicted targets (Table III), the majority of which were significantly decreased in grey compared with black hair follicles. Furthermore, among the miR-targeted DEGs, two genes (syndecan binding protein and KIT) are involved in pigment-associated pathways (26,27).

Another category of regulators affecting gene expression changes are TFs. Therefore, it was then examined whether the DEGs included any TFs and their potential targets. A total of 4 DEGs encoding TFs were identified: AE binding protein 1,
coiled coil domain-containing 17, runt-related transcription factor 3 (RUNX3) and SOX10, all of which were significantly decreased in grey hair follicles. Of note, two of these TFs, namely RUNX3 and SOX10, had potential targets among the other DEGs (Table IV), which also tended to be decreased in grey hair follicles (Fig. 2C), indicating an association between the downregulation of the TFs and their potential target genes in grey hair follicles. Transient receptor potential cation channel subfamily M member 1 (TRPM1), which encodes a permeable cation channel that is expressed in melanocytes and has a role in melanin synthesis, was among the downregulated genes regulated by these TFs (28,29). Of note, SOX10 have been proved to have the ability to drive the differentiation of melanocytes (30).
Hair graying, particularly premature hair graying, changes the appearance of affected individuals in a mostly undesired manner and has attracted the attention of researchers. To date, the underlying causes have remained largely elusive. In the present study, a genome-wide RNA-seq profiling analysis was performed using grey and black hair follicles from the same individuals with premature hair graying. It was revealed that pigment synthesis pathways were significantly impaired in grey vs. black hair, with a significantly decreased expression of multiple key genes crucial for the stability, trafficking and proliferation of melanocytes (31). The present results support the theory that premature graying may occur due to exhaustion of the melanocytes’ capability to produce hair pigmentation.

Previous studies have suggested that premature hair graying is associated with factors affecting melanogenesis, including nutritional deficiencies (32), insufficient neuroendocrine stimulation (33) and ionic signaling across melanosomes (34). The identified DEGs included those with similar functions with this regard, e.g. MCHR1, TRPM1 and SLCO4A2. MCHR1 acts as a receptor for melanin-concentrating hormone (35), TRPM1 regulates pigmentation at the plasma membrane level (29) and SLC45A2 modulates the melanosomal pH for optimal TYR activity required for melanogenesis (36). Of note, the present study identified that GJB1 was downregulated in grey vs. black hair follicles, which known to contribute to pigment transfer between melanocytes and neighboring keratinocytes (37).

While pigmentation-associated pathways are impaired in grey hair follicles, the underlying mechanisms of their inhibition/damage during hair greying remain to be elucidated. Of note, all of the 13 DEGs encoding miRs identified in the present study were significantly upregulated in grey hair follicles, indicating a substantial increase in the abundance of the corresponding mature miRs. Furthermore, the majority of the DEGs predicted to be targets of these miRs were significantly decreased in grey hair follicles, which is in line with the known mechanism that miRs reduce the expression of their target genes.

Another reason could be the decreased TFs in grey hair follicles. Among the four decreased TFs expressed in grey hair, two TFs (RUNX3 and SOX10) had potential target genes in other DEGs, which were also downregulated in grey hair, indicating that the downregulation of TFs were associated with their potential target genes in grey hair follicles. In addition, GJB1 was co-expressed with a crucial node, SOX10, in the PPI network. It has been reported that SOX10, in synergy with early growth response 2, may activate GJB1 in melanocytes, which may cause and alteration in melanogenesis (38,39).

Another point worth mentioning is that a relatively high fraction of DEGs was associated with the nervous system, including potassium voltage-gated channel subfamily Q member 2, basic helix-loop-helix family member e41, prostaglandin D2 synthase and centrosomal protein of 152 kDa, which are thought to be involved in controlling the circadian rhythm. Defects in these genes have been reported to be associated with a short sleep phenotype (40,41). It is widely accepted that nerve signaling defects, including a disturbed sleeping ability, may lead to hair graying, which
Table IV. Differentially expressed genes targeted by differential expression of genes encoding transcription factors (RUNX3 and sex-determining region Y box 10).

| Gene ID      | Gene       | FPKM in black hair | FPKM in white hair | Log2FC | P-value |
|--------------|------------|--------------------|--------------------|--------|---------|
| ENSG00000157404 | KIT       | 4.22               | 0.14               | -4.88  | 0.00005 |
| ENSG00000136040 | PLXNC1    | 6.36               | 0.46               | -3.79  | 0.00005 |
| ENSG00000104177 | MYEF2     | 1.29               | 0.17               | -2.92  | 0.00005 |
| ENSG00000020633 | RUNX3     | 1.48               | 0.26               | -2.49  | 0.00005 |
| ENSG00000145335 | SNCA      | 6.53               | 1.28               | -2.35  | 0.00005 |
| ENSG00000071575 | TRIB2     | 2.72               | 0.56               | -2.28  | 0.00005 |
| ENSG00000123095 | BHLHE41   | 5.33               | 1.2                | -2.16  | 0.01575 |
| ENSG00000130558 | OLFM1     | 1.27               | 0.33               | -1.94  | 0.00010 |
| ENSG00000137575 | SDCBP     | 10.59              | 2.79               | -1.93  | 0.00005 |
| ENSG00000048740 | CELF2     | 1.84               | 0.49               | -1.91  | 0.00005 |
| ENSG00000197283 | SYNGAP1   | 20.4               | 5.71               | -1.84  | 0.00240 |
| ENSG00000166173 | LARP6     | 1.39               | 0.4                | -1.8   | 0.01725 |
| ENSG00000026025 | VIM       | 26.35              | 7.8                | -1.76  | 0.00005 |
| ENSG00000204764 | RANBP17   | 1.35               | 0.43               | -1.65  | 0.02195 |
| ENSG00000115825 | PRKD3     | 6.13               | 2.31               | -1.41  | 0.00485 |
| ENSG00000082397 | EPB4L3    | 1.85               | 0.73               | -1.34  | 0.00015 |
| ENSG00000173482 | PTPRM     | 2.16               | 0.88               | -1.29  | 0.00870 |
| ENSG00000153823 | PTD1      | 1.42               | 0.59               | -1.26  | 0.00345 |
| ENSG00000115414 | FN1       | 1.68               | 0.72               | -1.23  | 0.01050 |
| ENSG00000185008 | ROBO2     | 1.5                | 0.65               | -1.2   | 0.04845 |
| ENSG00000086289 | EPDR1     | 3.04               | 1.42               | -1.09  | 0.00595 |
| ENSG00000116641 | DOCK7     | 11.49              | 5.51               | -1.06  | 0.00015 |
| ENSG00000249859 | PV1       | 10.11              | 20.29              | 1      | 0.01935 |
| ENSG00000135346 | CGA       | 14.88              | 32.89              | 1.14   | 0.00005 |
| ENSG00000137941 | TTLL7     | 1.77               | 4.44               | 1.32   | 0.00525 |
| ENSG00000105784 | RUNDC3B   | 0.44               | 1.41               | 1.7    | 0.00595 |
| ENSG00000113532 | ST8SIA4   | 0.33               | 1.21               | 1.88   | 0.01150 |
| ENSG00000151967 | SCHIP1    | 0.09               | 1.84               | 4.32   | 0.00015 |

miR, microRNA; FPKM, fragments per kilobase per million mapped reads; FC, fold change.
Figure 3. Protein-protein interaction network of the DEGs (TYR, TYRP1, MLANA, GPR143, OCA2, PMEL, SLC24A5 and SLC45A2). The network was generated using the STRING database by inputting the gene names of the DEGs between white hair and black hair follicles. DEG, differentially expressed gene.

Figure 4. Validation of 9 differentially expressed genes in white vs. black hair follicles. The abundance of target genes was normalized relative to the abundance of the β-actin gene. The values were calculated by using the $2^{-\Delta\Delta Cq}$ method. RNA was extracted from the gray and black hair of 10 patients. Each list indicates the log ratio of gray/black hair gene expression in each patient. A total of 10 columns present the log ratio. Nine genes are selected for quantitative analysis.
probably functions through its further effects on the nervous system and downstream pigmentation pathways (42). While it remains elusive whether and how the nervous system contributes to premature hair greying, the present results may indicate a novel aspect regarding the causes of hair greying. Since hair greying has been considered to be associated with aging of the hair follicle pigmentation system (43), the JenAge Ageing Factor Database (44) was searched for the DEGs in grey hair identified in the present study. A total of 5 genes were included in the online database, namely acetylcholine esterase, ATM, proprotein convertase subtilisin/kexin type 2, ubiquitin C-terminal hydrolase L1 and ventricular zone expressed PH domain containing 1, all of which had a decreased expression in white hair follicles, implying the decline of melanocyte-associated processes.

In conclusion, the present study was the first, to the best of our knowledge, to perform a genome-wide transcriptome profiling of human hair follicles affected by premature hair greying, which uncovered that damage of the melanin biosynthesis pathway was the direct cause of the decline hair pigmentation and the resulting hair greying. Furthermore, it was indicated that deregulated miRs and TFs, as well as neural disturbances, may be underlying causes. The present study provided multiple clues worthy of subsequent study to elucidate the mechanisms underlying canities and human premature hair greying. However, the present study is limited as only the processes occurring in subjects with premature hair greying were assessed, while the genetic predisposition to hair greying was not be determined. The genetic differences between subjects with premature hair greying and normal individuals should be assessed to identify the genes that are the primary cause of this condition.

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Availability of data and materials

The data used and/or analyzed during the present study are available from the corresponding author on reasonable request.

Authors’ contributions

DL, TN and HC conceived and designed the experiments. YB, XS and LY collected hair follicles, prepared RNA samples and constructed RNA-Seq libraries. GW analyzed transcriptome data. YB and LY performed qRT-PCR. YB and GW wrote the paper. All authors read and approved the final version of the manuscript.

Ethical approval and consent to participate

The present study was performed according to the declaration of Helsinki and was approved by the Research Ethics Committee at Fudan University (Shanghai, China). Informed consent was obtained from each participant prior to enrollment.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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