CAPN3, DCT, MLANA and TYRP1 are overexpressed in skin of vitiligo vulgaris Mexican patients

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Abstract. Vitiligo is a disorder causing skin depigmentation, in which several factors have been proposed for its pathogenesis: Environmental, genetic and biological aspects of melanocytes, even those of the surrounding keratinocytes. However, the lack of understanding of the mechanisms has complicated the task of predicting the development and progression. The present study used microarray analysis to characterize the transcriptional profile of skin from Vitiligo Vulgaris (VV) patients and the identified transcripts were validated using targeted high-throughput RNA sequencing in a broader set of patients. For microarrays, mRNA was taken from 20 skin biopsies of 10 patients with VV (pigmented and depigmented skin biopsy of each), and 5 biopsies of healthy subjects matched for age and sex were used as a control. A signature that identifies the expression pattern of 722 genes between depigmented vitiligo skin vs. healthy control, 1,108 between the pigmented skin of vitiligo vs. healthy controls and 1,927 between pigmented skin, depigmented vitiligo and healthy controls (P<0.05; false discovery rate, <0.1). When comparing the pigmented and depigmented skin of patients with vitiligo, which reflects the real difference between both skin types, 5 differentially expressed genes were identified and further validated in 45 additional VV patients by RNA sequencing. This analysis showed significantly higher RNA levels of calpain-3, dopachrome tautomerase, melan-A and tyrosinase-related protein-1 genes. The data revealed that the pigmented skin of vitiligo is already affected at the level of gene expression and that the main differences between pigmented and non-pigmented skin are explained by the expression of genes associated with pigment metabolism.

Introduction

Vitiligo is a skin disease characterized by the lack of pigmentation in the skin, it affects approximately 0.1 to 2% of the world population. However, its prevalence varies considerably among populations and ethnic groups: 0.14% in Russia, 1% in USA, and 2.5% in Japan (1), although the highest incidence has been described in Mexico (4%) and India (8.8%) (1,2). The most common clinical variant of vitiligo is vitiligo vulgaris (VV), in which the patient presents asymptomatic, well-circumscribed, milky-white macules involving one or multiple body regions or segments (3). The lack of pigmentation could be attributed to two main causes: a) the absence of melanocytes, which are dendritic cells derived from the neural crest that migrates to the epidermis and then to the hair follicle during embryogenesis, or b) the inability of these cells to produce and store melanin in melanosomes in the process of melanogenesis (4). In this context, the pathologic origin of vitiligo has not yet been fully understood. Several hypotheses and theories have been developed to explain these depigmentation processes (5-8).

Although melanocyte is responsible for the pigmentation process in vertebrates (9), the significance of the surrounding environment has been neglected, e.g., keratinocytes (8,10). Currently, it is known that the signaling mechanism that activates the route of melanogenesis is controlled by genes, whose products act as enzymes, structural proteins, transcriptional regulators, transporters, receptors and growth factors related

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to the melanogenesis process (11). Some of the hormones and products from the hypothalamic-pituitary-adrenal axis, and their respective receptors and negative regulators, trigger a nuclear signaling cascade that leads to the activation or repression of tyrosinase, a key enzyme in the pigmentation process, including the final amount of melanin produced (12,13). A major hormone regulator in melanin synthesis is the α-melanocyte-stimulating hormone (α-MSH), produced by pituitary gland, which interacts with a specific cell surface receptor [Melanocortin 1 Receptor (MC1R)] to stimulate melanin synthesis and other differentiated melanocyte functions (11).

Today, knowledge about genes and signaling pathways potentially involved in the development of vitiligo is increasing (14-17); for example, the expression profile of approximately sixteen thousand genes, in an in vitro culture of melanocytes obtained from five subjects with vitiligo, was analyzed using microarrays (17), describing five routes involved in the development of this disease: i) Development of melanocytes; ii) intracellular processing and vesicular trafficking of tyrosinase gene family protein; iii) packaging and transport of melanosomes; iv) cell adhesion; and v) processing and presentation of antigens (17).

In addition, previous reports of gene expression involved in the melanocortin system (14) and melanogenesis signaling pathways (15) showed modified expression levels of proopiomelanocortin (POMC), melanocortin 1 receptor (MC1R), melanocortin 4 receptor (MC4R), tyrosinase-related protein 1 (TYRP1) and dopachrome tautomerase (DCT), among other genes, in tissue affected by vitiligo when compared with healthy tissue samples from patients with vitiligo and normal skin from controls.

To date, there are few studies that involve the complete tissue analysis of patients with vitiligo. The identification of the expression profile of genes potentially involved in the development of the disease will be useful in understanding the molecular mechanism of its development to select potential therapeutic targets for its specific treatment.

In this study, we used microarray analysis to characterize the transcriptional profile of the skin of patients with VV and the identified transcripts were validated by the use of high-throughput RNA sequencing in a larger set of patients to determine the expression pattern that could play a role in the pathogenesis of vitiligo and the clinical types of this disease.

Materials and methods

Selection of participants. Fifty-five Northeastern Mexican patients from the states of Coahuila, Nuevo Leon, San Luis Potosi, Tamaulipas, and Zacatecas were recruited between November 2009 and May 2015 at the Dermatology Department of the University Hospital-UANL, in Monterrey, Nuevo Leon, Mexico. The Ethics and Research Committee of the Faculty of Medicine-UANL approved and registered the protocol and forms of informed consent under the code DE08-008 and DE13-001. After signing their informed consent, the patients were interviewed and evaluated to confirm the diagnosis of VV. Five healthy controls were also included for the Microarray analysis. None of VV patients had received any specific treatment in the previous six months to recruitment. The stage of VV (active AVV/stable SVV) was determined by intervals of time in the manifestation of new depigmented areas (stable vitiligo with lesional stability of >1 year), or enlargement of the already existing ones.

Sample collection and processing. Two skin biopsies of 4 mm were obtained from each patient with vitiligo. The first biopsy was obtained from the central part of the affected skin (called depigmented vitiligo skin) and the second from the healthy areas of the skin of patients with vitiligo (called pigmented vitiligo skin), generally 3 cm from the affected skin. Only a skin biopsy was taken from healthy control subjects (called control skin).

The skin tissue from the biopsies was immediately suspended in 5 volumes of RNA Later Solution (Ambion; Thermo Fisher Scientific, Inc., Waltham, MA, USA) after collection, and stored at 4 °C overnight. The next day, the supernatant was removed, and the samples were stored at -80 °C until the time for analysis.

Total RNA was isolated from the samples using the RNeasy fibrous tissue mini kit (Qiagen, Inc., Valencia, CA, USA) according to the manufacturer’s instructions and quantified using a NanoDrop® ND-8000 spectrophotometer (Thermo Fisher Scientific, Inc., Wilmington, DE, USA) and Qubit® RNA BR Assay kit (Thermo Fisher Scientific, Inc., Waltham, MA, USA). The quality/integrity of the extracted RNA was evaluated by an Experion Automated electrophoresis System (Bio-Rad Laboratories, Inc., Hercules, CA, USA).

Microarray assays. The biopsies taken from depigmented and pigmented skin from ten vitiligo patients (5 AVV and 5 SVV) and a skin biopsy of each of the five healthy controls were analyzed. For each sample, 100 ng of total RNA was amplified, purified, fragmentated and labeled using the GeneChip® IVT Express kit, (Affymetrix; Thermo Fisher Scientific, Inc.) according to the manufacturer’s instructions. 12.5 μg of fragmentated and labeled target were hybridized with Affymetrix GeneChip® Human Gene ul33 plus Array (Affymetrix; Thermo Fisher Scientific, Inc.), at 45 °C for 16 h, on a GeneChip® Hybridization Oven 640 (Affymetrix; Thermo Fisher Scientific, Inc.), according to the manufacturer’s recommendation. The hybridized arrays were washed and stained on a GeneChip® Fluidics Station 450, scanned on a GeneChip® Scanner 3000 7G (Affymetrix; Thermo Fisher Scientific, Inc.), and then CEL image data files were generated for each matrix.

Data analysis. CEL files were processed using the affy package in R (https://cran.r-project.org). Data were pre-processed using RMA and normalized using quantile-normalization (18). t-test and F-tests were used to determine a P-value for differentially expressed genes between groups. P-values were adjusted for multiple tests using the False Discovery Rate (FDR) approach (19). Principal component analysis (PCA) was performed in R.

Annotation and functional analysis. Further information about the genes was obtained from NetAffx™ Analysis Center (http://www.affymetrix.com) and NCBI databases (http://www.ncbi.nlm.nih.gov).
Functional annotation was performed by uploading the resulting gene list onto DAVID (20,21) (Database for Annotation, Visualization and Integrated Discovery, https://david.ncifcrf.gov/tools.jsp). Genes were mapped to KEGG-pathways and scored according to P-values (EASE Score, modified Fisher's exact test) and corrected for multiple testing according to the Benjamini-Hochberg False Discovery Rate correction provided by the DAVID tool.

**TruSeq Targeted RNA Expression analysis.** Skin biopsies from 45 vitiligo patients (23 AVV and 22 SVV) were analyzed by TruSeq Targeted RNA Expression (Illumina, Inc., San Diego, CA, USA). RNA was obtained and cDNA was prepared with ProtoScript® Reverse Transcriptase (New England Biolabs, Ipswich, MA, USA) according to the manufacturer’s instructions. RNA expression analysis was performed on an Illumina MiSeq with MiSeq® Reagent Kit v3 (150 cycle) and TruSeq® Targeted RNA Custom Panel Kit designed to detect the expression profile of the calpain 3 (CAPN3), dopachrome tautomerase (DCT), glycerol-3-phosphate dehydrogenase 1 (GPD1), melan-A (MLANA) and tyrosinase-related protein 1 (TYRP1). RNA-Seq data were analyzed within Base Space (https://basespace.illumina.com) using the tool TruSeq Targeted RNA. Count data were exported and normalized per sample by equalizing the total number of counts (multiplying by the average of the total counts per sample then dividing by the accumulated counts per sample).

**Results**

**Microarray analysis.** In the analysis performed on vitiligo samples using microarray analysis, we first compared whether the unaffected skin holds a higher similarity to healthy controls than to the affected skin of the patients; one of the pigmented skin samples from a vitiligo patient was discarded from the microarray analyses as it lacked sufficient quality to be used. A PCA and hierarchical clustering shows that the unaffected skin is more similar to vitiligo samples than to healthy controls (Fig. 1), suggesting that pigmented skin in vitiligo patients is already affected regardless of its clinical evidence.

Further analysis comparing the healthy controls vs. skin samples from vitiligo patients (de/pigmented tissue) confirmed the existence of a large number of hits (1,927 probes at FDR <0.1) (Fig. 2).

Then we compared different combinations of groups at FDR <0.1, showing that there are differences between the three groups, with altered expression patterns (over and under expressed) when analyzing the more than 25,000 genetic targets present in Affymetrix Gene Chip® Human Gene u133 plus Array. Further, we observed several changes when comparing the healthy controls and patients independent of sample type (pigmented or depigmented skin of patients with vitiligo) that reached the 1,927 probes (Table I). When comparing the pigmented tissue of vitiligo patients against healthy controls, the difference decreases to 1,108 (Table I). Surprisingly, the number decreases further to 722 when comparing the depigmented vitiligo samples to healthy controls (Table I). Finally, the differences are modest when comparing the vitiligo depigmented samples against pigmented asymptomatic tissue, consistent with the initial observation that the asymptomatic tissue is already compromised. Therefore, we focused on these minor differences between vitiligo samples and their healthy counterparts, observing the differences in the important components of pigmentation (Fig. 3). Moreover, we noted that the expression profile of these genes were similar to those from healthy controls. On the other hand, no relationship was found between expression pattern variations, gender, age, or type of vitiligo.

**Functional Annotation of observed differences.** Using the DAVID functional annotation tool, the genes identified in
involved in skin pigmentation: DCT, MLANA and TYRP1. In addition, CAPN3 was also significant, although at a lower level. In addition to microarray analysis, no relationship was identified between the variants in the patterns of expression, gender, age, or type of vitiligo. Moreover, in the case of the GAPD1 gene, no significant difference in the expression pattern between asymptomatic skin and vitiligo lesions was found by NGS when compared to previous observations using microarray analysis (Fig. 5).

Discussion

Most vitiligo studies conducted in the Mexican population are focused in demographics and clinical characteristics. To date, the molecular etiology and genetic factors interacting in the development of vitiligo within this population have not been considered.

Up to date, there has been only two expression studies of vitiligo using microarray technology. In the first study, melanocyte culture in vitro from 5 samples of VV patients were analyzed (17); in addition to the small number of samples, it did not reflect the contribution of the skin’s environment as it only represents the alterations of a single cellular type (melanocytes); further, the expression pattern could be modified under cell culture conditions. This report found altered an expression pattern of genes involved in the development and function of melanocytes, proposing an autoimmune response as a secondary event caused by the abnormal function of melanocytes (17). In our study, we also observed some similarities in the expression pattern of genes involved in pigment synthesis, melanosomes, granule pigment, cytoplasmic vesicles (possibly melanosomes), plus redox reaction processes related to diverse functions, including cell-cell recognition, cell-surface receptors, muscle structure, and the immune system. However, our results reflect the expression profile of the whole skin.

In a second report, an expression study using the Smyth line from chicken, an avian model for human autoimmune vitiligo, the microarray analysis results support the multifactorial etiology of vitiligo, where inflammatory/innate immune activity and oxidative stress, and the adaptive immune response play a predominant role in melanocyte loss. The microarray analysis results provided comprehensive information at transcriptome level, supporting the multifactorial etiology of vitiligo, where, along with the apparent inflammatory/innate immune activity and oxidative stress, the adaptive immune response plays a predominant role in melanocyte loss (22).

In the biological context of the skin, the melanocyte represents less than ten percent of epidermal cells (23,24). The functional clustering analysis made using the DAVID database points towards the involvement of an altered functional group composed by genes involved in diverse biological processes, such as transcription and transcription regulation, transcriptional repression, alternative splicing, and keratin filament. The observation that melanocytes are the main cell type affected in vitiligo does not rule out the participation of other cellular components of the skin in the development of this disease, as supported by the major affected pathways of pigment synthesis, packaging and transport of pigment, when the depigmented vitiligo tissue was compared with the pigmented skin of
For this reason, and considering that keratinocytes are the predominant cell type in this tissue (25), their role could involve an important function in skin pigmentation (10,26). Even when the differences in expression patterns between a) pigmented skin and b) affected skin vs. control skin in the same metabolic routes are affected, in the first case a greater number of genes are involved (Table 1), these could be explained by compensatory mechanisms (14,27) in which the skin of the

| Comparison/N | Biological altered pathways |
|--------------|----------------------------|
| Vitiligo depigmented skin vs. vitiligo pigmented skin vs. controls (N=1,927) | Alternative splicing/Splice variant DNA-binding/Nucleic acid binding/Zinc-finger/Transcription factor and transcription regulation/Krueppel-associated box Metal-binding Phosphoprotein Krueppel-associated box Coiled coil |
| Vitiligo pigmented skin vs. controls (N=1,108) | Alternative splicing/Splice variant Intracellular Krueppel-associated box Coiled coil |
| Vitiligo depigmented skin vs. controls (N=722) | Keratin filament High sulphur keratin-associated protein Splice variant Krueppel-associated box |
| Vitiligo depigmented skin vs. vitiligo pigmented skin (N=6) | Melanosome/melanosome membrane Uncharacterised domain, di-copper centre Tyrosinase/Tyrosine metabolism Topological domain: Lumenal, melanosome Melanin biosynthesis/Melanogenesis |

(N) Altered genetic target from of 25,000 targets present in Affymetrix GeneChip® Human Gene u133 plus Array.

Figure 3. Biological processes involved in skin pigmentation and development of vitiligo. On the left are presented routes in normal skin pigmentation process; on the right are represented the major routes detected as altered in skin of vitiligo subjects.
The vitiligo patient tends to stimulate pigmentation by regulating the expression factors involved in pigmentation, increasing or decreasing their expression to maintain skin homeostasis.

After analyzing the genes showing a different expression pattern, a group of 5 genes, involved in the melanogenic routes, intracellular cysteine proteases, and oxidative stress, were selected for validation by TruSeq Targeted RNA Expression analysis. The validation assays showed a differential expression tendency in four genes: CAPN3, DCT, MLANA and TYRP1. However, GPD1 was not found differentially expressed by NGS. These results are consistent with Regazzetti et al. (28), whom validated a group of genes in skin biopsies from patients with vitiligo, finding low expression levels of MLANA, DCT, and TYRP1 in vitiligo skin lesions when compared to skin around the injured and uninjured (pigmented/asymptomatic) tissue.

The CAPN3 gene was under expressed in the skin of patients with vitiligo; previously, Stromberg et al reported that this gene was up regulated in melanocyte cultures obtained from vitiligo lesions (17). This gene encodes a large subunit of neutral calpain of protease 3 activated by muscle calcium, a heterodimer consisting of a large subunit and a small subunit is a major intracellular protease that is ubiquitously expressed in human tissues and other and other tissue-specific isoforms. Its function on the skin is not clear, but it has been observed that CAPN3 variants can play a pro-apoptotic role in melanoma cells and its down regulation, as observed in highly aggressive melanomas, could contribute to their progression (29). In melanocytic cells, CAPN3 expression may be regulated by MITF (30) which regulates a broad variety of genes whose functions range from pigment production (such as DCT, MC1R, MLANA, TYR and TYP1) to the cell-cycle regulation, migration and survival; moreover, MITF-mediated up regulation of CAPN3 has been reported in melanoma (30).

The GPD1 gene encodes a member of the NAD-dependent glycerol-3-phosphate dehydrogenase protein family, playing a critical role in carbohydrate and lipid metabolism (31). Along with GPD2, the mitochondrial isoform, it constitutes a glycerol phosphate shuttle that facilitates the transfer of reducing equivalents from the cytosol to the mitochondria, playing a crucial role in osmoregulation and redox balance (32). In our study, the microarray analysis showed the under expression of this gene in tissue from patients with vitiligo lesions. As a result of expression profile validation, made by RNA-Seq in a number of samples, no significant differences were observed between the expression profiles of affected pigmented skin in vitiligo patients. However, we observed a higher expression in some of the samples that possibly participate in the oxidative stress response experienced by the skin cells of these patients, a mechanism that has been described in this disease. So far, the participation of this gene in vitiligo has not been explored.

In addition, NGS provides a better approach to gene expression profile analysis, as this technology enables high resolution research for all of the RNA present in a sample, including mRNA sequence and abundance (33). Its usefulness has been compared with microarray technology and Real-Time PCR in the analysis of expression profiles (34-36); therefore, it has been used in transcriptome analysis in clinical research (36-39).

The analysis of expression by directed sequencing of RNA (RNAseq) has been helpful in the efficient evaluation of expression profiles of multiple genes in human pathological and non-pathological conditions (40-43). In particular, RNAseq using NGS, not only shows the expression levels of a transcript, but the presence of isoforms in the samples analyzed (44). In patients affected by dermatological diseases such as psoriasis, systemic lupus erythematosus (45-47), and vitiligo, these tools are an excellent alternative in identifying modified gene expression patterns in key routes of skin.
homeostasis, pigmentation, and cell survival at a low cost and reduced time in comparison to RT-qPCR.

In conclusion, we found alterations in the expression pattern of degeminated vitiligo lesions when compared to pigmented asymptomatic skin of vitiligo patients and against healthy control samples. These results reflect that even the clinically unaffected (pigmented) skin shows an impaired melanogenesis process in a patient with vitiligo; the observed changes are related to processes regulating gene expression and splicing that eventually will end up in skin depigmentation by altering the levels of proteins involved in this process. Upregulated expression profiles detected by microarray analysis for CAPN3, DCT, MLAN-A and TYRP1 in VV patients was validated using NGS TruSeq Targeted RNA Expression analysis, and whose altered functions on pigment production, and possibly in melanocyte cell survival, affect skin pigmentation, thus suggesting that these tools are an excellent alternative in identifying the expression pattern of genes involved in the development of this disease. On the other hand, we did not find any differences in the gene expression profiles of active or stable VV.

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