The genetics and epidemiology of generalized vitiligo

Generalized vitiligo (GV) is the most common depigmentation disease, in which white spots of skin and overlying hair result from loss of melanocytes from the involved regions. GV is a complex disease involving both genetic predisposition and unknown environmental triggers. Whereas various pathogenetic mechanisms have been suggested, most evidence supports an autoimmune basis for this disease. Recently, three different genome-wide association studies of GV have been reported, identifying a total of 17 confirmed GV susceptibility loci. Almost all of these genes encode immunoregulatory proteins, together highlighting pathways by which melanocytes might be recognized and killed. Moreover, the biological interaction between two of these GV susceptibility genes, HLA-A and TYR (encoding tyrosinase), points to an apparent inverse relationship between susceptibility to GV versus malignant melanoma, suggesting that GV may result, in part, from dysregulation of normal processes of immune surveillance against melanoma.

Vitiligo susceptibility genes

Candidate gene association studies and gene expression analyses have produced a long list of GV candidate genes, of which only HLA (human leukocyte antigen) and PTPN22 (protein tyrosine phosphatase, non-receptor type 22) have had consistent support from multiple studies; most of the rest are likely to represent false positives [3,6]. Two additional genes, NALP1 (now NLRP1; NLR family, pyrin domain containing 1) and XBPI (X-box binding protein 1), were first mapped by unbiased genome-wide linkage analyses and subsequently were identified by positionally targeted genetic association studies [7,8]; both of these genes have subsequently been replicated by multiple studies [3,6]. These four confirmed GV susceptibility loci - HLA, PTPN22, NLRP1 and XBPI - all encode important immunoregulatory proteins, lending support to the autoimmune hypothesis of GV pathogenesis.

Recently, three different genome-wide association studies (GWASs) of GV have been reported: two from...
European-derived white populations [9-11] and one from China [12]. Together, these studies identified a total of 17 confirmed GV susceptibility loci, yielding major insights into pathways of disease pathogenesis and overall strongly supporting an autoimmune basis for typical GV. The first GWAS, of a founder population in an isolated Romanian village with a high prevalence of GV and other autoimmune diseases [9], detected association at chromosome 6qter near IDDM8, which is a type 1 diabetes–rheumatoid arthritis locus in the vicinity of SMOC2 (SPARC-related modular calcium binding 2). The second GWAS [10,11], also carried out in European-derived white individuals, identified a total of 13 susceptibility loci for GV, including HLA class I (specifically, HLA-A*0201), HLA class II, PTPN22, RERE (arginine–glutamic acid dipeptide repeats), FOXP1 (forkhead box P1), LPP (LIM domain containing preferred translocation partner in lipoma), CCR6 (chemokine (C-C motif) receptor 6), IL2RA (interleukin 2 receptor, α), TYR, GZMB (granzyme B), NLRP1, UBASH3A (ubiquitin-associated and SH3 domain containing A) and CIQTNF6 (C1q and tumor necrosis factor related protein 6). Moreover, subsequent re-analysis of this genome-wide dataset to specifically test association of 33 biological candidate genes previously implicated in GV identified three additional GV susceptibility loci: TSLP (thymic stromal lymphopoietin), XBP1 and FOXP3 (forkhead box P3) [6].

The third GWA study [12], carried out in a Chinese population, also identified GV susceptibility loci in the HLA class I and class III regions, and likewise detected association with CCR6. Moreover, the GV-associated SNP in the CCR6 region is the same in European-derived white individuals and Chinese people, suggesting that these two populations may share a single causal allele, and is located only 1.44 Mb from the previous GV-associated SNP in the SMOC2 region [9], suggesting the possibility that these two signals might be related. Virtually all of the confirmed GV susceptibility loci encode known immunoregulatory proteins, and many have been associated with genetic susceptibility to other autoimmune diseases that are epidemiologically linked to GV (Figure 2).

The one exception among the susceptibility loci associated with GV is TYR, which encodes tyrosinase, the key enzyme of melanin biosynthesis in melanocytes. However, in GV even TYR may act primarily to modulate recognition of the melanocyte target cell by the immune system. Beyond its role in pigmentation, tyrosinase is the major autoantigen in GV. In this regard, GV is thus analogous to type 1 diabetes and autoimmune thyroid disease, in that genetic susceptibility to disease involves genes that encode key specialized intracellular components of the autoimmune target cell types and that constitute major autoantigens for the corresponding disease (GV: TYR, tyrosinase; type 1 diabetes: INS, insulin; autoimmune thyroid disease: TG, thyroglobulin). For GV, the causal TYR susceptibility variant appears to be the major (Arg) allele of rs1126809, a common non-synonymous (Arg402Gln) polymorphism that has a minor allele frequency of 0.22 to 0.40 in European-derived white individuals. This polymorphism is rare in other populations, which is why it was not detected in the Chinese GWAS, even though TYR may well play a role in GV pathogenesis in all populations. In contrast, the minor (Gln) allele, which is protective with respect to GV, is associated with susceptibility to malignant melanoma in European-derived white individuals [13,14]. Thus, from the standpoint of genetic susceptibility, the TYR Arg402Gln polymorphism represents an inverse relationship between GV and malignant melanoma.

Much of the biology that is likely to underlie this inverse relationship is already known, largely from extensive studies on melanoma patients, in whom GV may develop during the course of treatment and is an auspicious prognostic sign. Tyrosinase is a major antigen
presented to the immune system on the surface of melanocytes and melanoma cells by HLA class I molecules, principally \( HLA-A^\ast0201 \), which itself is a major GV risk allele. Indeed, \( HLA-A^\ast0201 \) and \( TYR 402\text{Arg} \) exhibit significant genetic interaction in promoting GV susceptibility [10], reflecting a corresponding biological interaction. One of the important class-restricted epitopes presented by \( HLA-A^\ast0201 \) is a specific modified tyrosinase peptide: YMDGTMSQV [15]. However, the \( TYR 402\text{Gln} \) variant results in an unstable polypeptide that is retained in the endoplasmic reticulum and degraded, thereby reducing the amount of tyrosinase peptide available for presentation on the cell surface. Moreover, presentation of this tyrosinase peptide by \( HLA-A^\ast0201 \) requires the posttranslational modification of residue 371Asn to Asp [15], via a mechanism that is probably inefficient in the \( TYR 402\text{Gln} \) polypeptide. Tyrosinase is thus an important signal by which the immune system recognizes melanocytes, and tyrosinase-402Arg is likely to make a greater contribution than tyrosinase-402Gly to immune surveillance (and thus protection) against melanoma cells. Thus, at least four of the GV genome-wide association signals - \( HLA \) class I, \( TYR \) 402Arg, \( IL2RA \) and \( GZMB \) - may be part of a pathway by which CTLs recognize and ultimately kill melanocytes in patients with GV, and perhaps also protect against incipient melanomas.

These observations open several doors to future studies. In GV, it has been known for some time that patients have circulating skin-homing melanocyte-specific CTLs [16]. However, the molecular epitopes recognized by these melanocyte-specific CTLs have not yet been identified. At least in GV patients carrying \( HLA-A^\ast0201 \), one of these CTL epitopes might be the modified tyrosinase peptide YMDGTMSQV. Furthermore, it is not yet known whether susceptibility to GV is generic with respect to \( HLA-A^\ast0201 \) (which is quite common), or whether it is particular to specific subtypes, and whether those subtypes conversely confer the greatest protection against melanoma.

**Conclusions and perspectives**

Recent GWASs have yielded substantial progress in identifying genes involved in risk of GV, with 17 loci now confirmed (\( HLA \) class I, \( HLA \) class II, \( HLA \) class III, \( PTPN22 \), \( RERE \), \( FOXP1 \), \( LPP \), \( TSLP \), \( CCR6 \), \( IL2RA \), \( TYR \), \( GZMB \), \( NLRP1 \), \( UBASH3A \), \( XBP1 \), \( C1QTNF6 \), and \( FOXP3 \)): 16 in European-derived white individuals and...
four in Chinese people, and, for a few genes, in both. Nevertheless, the 16 loci identified in European-derived white individuals together account for only 10% of the total genetic risk of GV in that group, indicating that additional loci probably remain to be discovered, with a few common and perhaps numerous rare variants accounting for disease risk at each locus. Essentially all of the confirmed GV susceptibility genes regulate function of the immune system, and many have also been associated with other autoimmune diseases, highlighting shared pathways of autoimmune susceptibility among these diseases.

Furthermore, findings for both HLA-A and TYR suggest an inverse relationship between susceptibility to GV and susceptibility to malignant melanoma, with genetic interaction that reflects underlying biochemical and functional interaction between the corresponding proteins. The overall picture indicates that genetic variation at HLA-A*0201 and TYR interacts to modulate immune surveillance against malignant melanoma, with heightened surveillance predisposing to GV and protecting against melanoma, and reduced surveillance protecting against GV but predisposing to melanoma. This biological relationship may also explain the frequent occurrence of GV in patients treated for melanoma, in whom development of this autoimmune phenotype constitutes a relatively favorable prognostic sign.

While these discoveries underscore the autoimmune nature of GV, they do not offer specific clues as to the environmental triggers that may initiate the autoimmune response. GV is a disease of the skin, the organ that is the first point of contact for one’s interaction with the external environment, and which is highly accessible for analysis of that interaction. Furthermore, because GV is so visually evident, it is usually diagnosed early. Thus, identification of GV susceptibility genes may enable identification of individuals at high genetic risk, enabling relatively direct analysis of potentially causal gene-environment interactions, both retrospectively in patients with relatively recent disease onset, and prospectively in individuals who are at high genetic risk.

Abbreviations
CTLT, cytotoxic T lymphocyte; GV, generalized vitiligo; GWAS, genome-wide association study; SNP, single-nucleotide polymorphism.

Competing interests
The author declares that he has no competing interests.

Author's information
Dr Richard A Spritz is Professor of Pediatrics and Director of the Human Medical Genetics Program at the University of Colorado School of Medicine. The work of Dr Spritz in the field of human genetics began in the ‘pre-molecular’ era, and in the mid-1970s he took part in the first cloning of human genes and identification of the first human disease gene mutation. Over the past 30 years, Dr Spritz and his colleagues have studied the genes involved in causing many different human diseases, including hemoglobin disorders, albinism and other skin diseases, autoimmune diseases such as vitiligo and thyroid disease, and cleft lip/palate, and he has published over 200 scientific papers on these investigations. Most recently, Dr Spritz led an international team that carried out the first genome-wide association study for generalized vitiligo.

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