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Risk Factors and Disease Mechanisms in Myositis

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Abstract

Autoimmune diseases develop as a result of chronic inflammation owing to interactions between genes and the environment. However, the mechanisms by which autoimmune diseases evolve remain poorly understood. Newly discovered risk factors and pathogenic processes in idiopathic inflammatory myopathy (IIM) phenotypes have illuminated innovative approaches for understanding these diseases. The HLA 8.1 ancestral haplotype is a key risk factor for major IIM phenotypes in white populations, and genetic risk variants for other autoimmune diseases have been identified as IIM risk factors. Environmental risk factors are less studied but might include viruses, bacteria, ultraviolet radiation, smoking, occupational and perinatal exposures and a growing list of drugs, biologics, and dietary supplements. Disease mechanisms vary by phenotype, with evidence for shared innate and adaptive immune and metabolic pathways in some phenotypes but unique pathways in others. The heterogeneity and rarity of the IIMs make advancements in diagnosis and treatment cumbersome. Novel approaches, better-defined phenotypes, and international, multidisciplinary consensus have contributed to progress, and hopefully these methods can eventually enable therapeutic intervention before the onset or major progression of disease. In the future, preemptive strategies to IIM management might be possible.

Introduction

The pathomechanisms of most immune-mediated diseases relate to chronic organ inflammation that can be caused by specific interactions between genetic and environmental risk factors. Immune activation in such diseases often involves both innate and adaptive

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mechanisms, as well as other non-immune mechanisms; however, the details and the interactions of different pathways are usually not clear. The rarity and heterogeneity of the idiopathic inflammatory myopathies (IIM), a group of systemic autoimmune diseases which include polymyositis, dermatomyositis, necrotizing myopathy, myositis that is found in conjunction with other autoimmune diseases, called overlap syndromes, and inclusion body myositis (IBM) (Box 1), have hampered our understanding of their risk factors and pathogenesis. Nonetheless, considerable progress has been made in this area over the last decade (Figure 1).

In this Review, we describe the major advances from the past few years in our understanding of the genetic risk factors, associated environmental exposures, and immune mechanisms and non-immune-mediated mechanisms in IIM phenotypes. Given space limitations, we do not cover many historical features or the details and biologic differences among the phenotypes. The classification, outcome assessments, detailed autoantibody developments and therapies in the IIMs are reviewed in accompanying articles.

Genetic risk factors

Genetic epidemiology of IIM

Owing to the rarity of IIMs, there are few reports of familial occurrence^{1,2}; hence, the heritability of IIMs is unknown. This caveat contrasts with other autoimmune disorders, such as rheumatoid arthritis or type I diabetes, which have reported heritabilities of 66%-88%^{3,4}. In comparison with healthy individuals, higher prevalences of autoimmune disease, such as systemic lupus erythematosus, autoimmune thyroid disease, or type 1 diabetes, have been reported concurrently in patients with IIM, as well as in first-degree relatives of both adult and juvenile patients with IIM, on the basis of data from a relatively small number of patients^{5,6}. Conversely, a nationwide study in Taiwan that investigated co-aggregation of autoimmune disease in families with systemic lupus erythematosus and systemic sclerosis, identified higher relative risk of IIM than that of the general population^{7,8}, and a national study in China suggested an increased risk of IIM and certain other autoimmune diseases in relatives of patients with systemic sclerosis⁸. These findings of aggregation of autoimmune diseases within families of patients with IIM suggest that shared genetic and/or environmental factors might contribute to disease risk. However, large, well-powered, epidemiological studies are needed to robustly evaluate these data. Notably, in 2015, a systematic review demonstrated that the reported incidence and prevalence rates for IIMs have increased over time⁹. Although this finding might reflect a true increase in disease burden, this increase could also be due to a wider recognition or more accurate recording of disease than in the past.

HLA loci associated with IIM

Genome-wide single-nucleotide polymorphism (SNP) association studies in adult and juvenile individuals of European ancestry who have dermatomyositis or polymyositis identified the strongest disease associations within the MHC region on chromosome $6^{10,11}$, one of the most complex regions of the genome, which has a high concentration of genes encoding proteins with immunological functions. The development of the so-called

Immunochip (Illumina, USA), a cost-effective genotyping array that includes ~200,000 genetic variants associated with autoimmune diseases, combined with accurate imputation of HLA alleles and amino acids from SNP data¹², greatly improved our ability to dissect HLA associations. The Myositis Genetics Consortium conducted the largest genetic study in IIM, which included 2,566 patients with IIM from 14 countries. The study demonstrated the strongest disease association with alleles of the 8.1 ancestral haplotype—*HLA-DRB1*03:01* and *HLA-B*08:01* in polymyositis and dermatomyositis, respectively—whilst conditional analysis revealed that multiple variants on this haplotype might contribute independently to disease risk¹³. In IBM, *HLA-DRB1*03:01, HLA-DRB1*01:01,* and *HLA-DRB1*13:01* were independently associated with disease¹⁴; the latter two alleles are uniquely associated with IBM. Different risk factors are seen in other ethnic groups, including HLA-*DRB1*07,* which is associated with dermatomyositis in Chinese populations¹⁶.

Specific amino acid associations in the HLA region, such as position 57 of HLA-DQB1, position 77 of HLA-DRB1*03:01, and positions 26 and 11 of HLA-DRB1*03:01, differentiate dermatomyositis, polymyositis, and IBM, respectively^{13,14}, and suggest different predominating pathophysiology in different clinical subgroups. Conversely, although HLA allele associations differ in non-white populations, the amino acid associations might be consistent across different populations. Amino acid sequence variations might alter the structure of the HLA molecule's peptide-binding groove and thereby increase disease susceptibility by influencing antigen repertoires and the affinity of peptides presented to the immune system. If so, computational modelling of predicted antigen-HLA binding to identify the immunogenic peptides could help to determine disease mechanisms.

The finding that different HLA alleles have been associated with various myositis-specific autoantibody (MSA)-defined subgroups¹⁷ agrees with the finding that many MSAs are mutually exclusive. In general, HLA risk alleles are more strongly associated with MSAdefined subgroups than clinically-defined subgroups (which are less homogenous than the MSA-defined subgroups) despite smaller sample sizes in the MSA groups. For example, the presence of anti-histidyl-tRNA synthetase (anti-Jo1) autoantibodies in IIM is strongly associated with multiple alleles of the 8.1 ancestral haplotype, including HLA-B*08:01, DQB1*02:01, and DRB1*03:01, particularly when multiple alleles are considered together as a haplotype¹¹. The presence of other autoantibodies, including autoantibodies to Mi-2 (also known as chromodomain-helicase-DNA-binding proteins), SUMO-activating enzyme (SAE), melanoma differentiation-associated gene 5 (MDA5), signal recognition particle (SRP), transcription intermediary factor 1 (TIF1) and anti-PL-7, have been associated with specific HLA alleles^{17,18}. Although most HLA associations are the same in adult and juvenile IIM phenotypes, HLA associations can occasionally differ between adult and juvenile patients, as is the case with anti-3-hydroxy-3-methylglutaryl-coenzyme A reductase (HMGCR) autoantibodies, with associations of HLA-DRB1*11:01 and HLA-DRB1*07:01 for adult and juvenile patients with IIM, respectively^{19,20}.

Non-HLA loci associated with IIM

Several non-HLA loci have been associated with IIM (Table 1), including *PTPN22*^{13,21}; this locus was associated with polymyositis (reaching genome-wide significance, defined as $p<5\times10^{-8}$), but not with adult or juvenile dermatomyositis¹³, again suggesting that different clinical subgroups have different pathophysiologies. Other genes, including *STAT4*, *TRAF6*, and *UBE2L3* in IIM^{13,22}; *PLCL1* and *BLK* in dermatomyositis^{10,23}; and *CCR5* in IBM¹⁴, have been associated with disease, reaching a suggestive level of significance (defined as $p<2.25\times10^{-5}$). Complement 4A (C4A) deficiency, owing to copy number variation, has been linked to an increased risk of developing juvenile dermatomyositis, although the strongest risk factor identified in the study was attributable to the presence of *HLA-DRB1*03:01* along with C4A deficiency²⁴. In IBM, sequencing of candidate genes involved in related neuromuscular or neurodegenerative diseases^{25,26} and whole-exome sequencing of proteins overrepresented in the skeletal muscle rimmed vacuoles—defined as a space within the cytoplasm of a muscle cell with a purplish staining rim on trichrome staining—of patients with IBM²⁷ identified rare variants in *VCP*, *SQSTM1*, and *FYCO1* associated with disease, suggesting impaired autophagy as a mechanism of IBM pathogenesis.

In contrast to the more common immune-mediated diseases, where extensive meta-analyses have been conducted, a relatively small number of genetic risk variants has been identified for IIM. This small number likely reflects the sample sizes of patients with IIM, as well as the marked heterogeneity of these complex diseases, and highlights the importance of collaborative endeavours. Most large genetic studies of IIMs have focused on populations of European ancestry. Further large-scale studies are required to establish whether variants, pathways, and gene–environment interactions are shared across different ethnic groups.

Pathways implicated in IIM pathogenesis

By identifying the genes associated with IIM, studies can focus on the molecular pathways involved and thereby improve our understanding of IIM pathogenesis. The strong association between IIM subsets and *HLA-DR* and *HLA-DQ* genes supports a role for the adaptive immune system in the pathogenesis of IIM, as a key role of HLA class molecules is to present antigens to T cells. The roles of other associated variants have been investigated by functional annotation, for example, analysing the effects of coding variants on the translation of the encoded protein and/or regulatory effects on gene expression through expression quantitative trait loci (eQTL) analysis. Identification of eQTLs might also help to identify functionally relevant cell types through immune cell–specific eQTLs that affect spatial and temporal gene expression, the cellular response to stimulation, and/or the magnitude of the response. Associated genes (Table 1) implicate both the innate and adaptive immune responses through, for example, the roles of *PTPN22* and *STAT4* in the T cell receptor pathway, or *BLK, UBE2L3*, and *TRAF6* in B cells and the nuclear factor κ B (NF- κ B) signalling pathway. In IBM, specific genes implicate both inflammatory and degenerative changes, including mitochondrial abnormalities, in disease pathogenesis.

Despite the small contribution of identified genetic variants to clinical phenotypes, drugs targeting the pathways affected by genetic variations might be disproportionately effective. In IIM, the application of drugs repurposed from other diseases will probably become more

important as our understanding of disease mechanisms evolve. Refinements in defining the genetic factors that drive different phenotypes will be important in clinical decision-making for early and effective diagnosis, classification, and therapeutic management, by targeting therapy to patients most likely to respond.

Missing heritability

The pathogenesis of IIM cannot be explained solely by genetic risk factors. Many of the variants identified have a relatively small effect on disease risk individually, and only 5.5%–16% of the phenotypic variance can be explained by genetic variants identified from Immunochip studies²⁸. Following accepted models of genetic architecture, rare variants, including single nucleotide and copy number variants, are probably involved in rare diseases such as the IIMs but have not yet been extensively investigated. This postulate is illustrated by Mendelian forms of monogenic juvenile-onset disorders that share clinical and immunologic features with juvenile dermatomyositis, such as chronic atypical neutrophilic dermatosis with lipodystrophy and elevated temperature (CANDLE) syndrome²⁹.

Environmental risk factors

Although a number of genes have been associated with the IIMs, the physiologic effect of these genes might depend on their activation or modification by environmental factors. Compared with genetic studies, however, fewer investigations have explored the role of environmental exposures in the development of IIM. Thus, the field is decades behind other areas, such as oncology, which has identified modifiable risk and protective factors for different cancers; by contrast, in many cases, only a single or small number of studies have identified specific environmental exposures associated with particular IIM phenotypes.

Multiple lines of evidence suggest that autoimmune diseases have an environmental component: the concordance rates for autoimmune diseases in monozygotic twins is much less than $50\%^{30,31}$; there are strong temporal associations between certain exposures (infectious agents and drugs in particular) and the subsequent development of some autoimmune diseases^{32–34}; in some individuals, disease improves after removing a suspected environmental agent (dechallenge) and/or worsens or reoccurs after re-exposure to the suspected agent (rechallenge)^{34–36}; the incidence of many autoimmune diseases has increased over time^{37,38}; there are seasonal and geographic variations in disease onset and in birth dates of individuals who have developed an autoimmune disease^{34,39–42}; data from relevant animal models have demonstrated the plausibility of multiple environmental agents potentially triggering autoimmune disease^{43–45}; the major genetic risk factors for autoimmunity are polymorphic genes that regulate responses to environmental agents^{46,47}; variations in the human immune system are largely driven by nonheritable influences⁴⁸; and associations between specific exposures and autoimmune diseases have been documented in large epidemiologic studies^{49,50}.

In addition to their role as possible initiators of autoimmunity, environmental factors might alter the rate of progression to clinical disease, the specific manifestations of disease expression, and/or the course of illness^{34,51}. After certain disease-initiating exposures, decades can pass before autoimmune disease manifests, and, as for many cancers, there can

be a progression of stages from autoimmunity to preclinical disease with immune alterations to classifiable autoimmune disease⁵². A panel of international investigators convened by the U.S. National Institute of Environmental Health Sciences in 2012 defined a number of well-supported associations between environmental exposures and autoimmune diseases⁵⁰ and proposed criteria to define environmentally associated autoimmune diseases in clinical care and epidemiologic settings³⁶. Thus, there are many possible ways in which environmental agents might influence individuals over the lifetime to result in perturbations that could eventually result in the autoimmune phenotypes we recognize today. However, it is critical to develop consensus on how to best recognize and diagnose environmentally associated autoimmunity.

Specific environmental agents of concern

As with other autoimmune diseases, many environmental studies in IIM have been based on animal models, case reports, and/or case series, and which suggest environmental risk factors that might vary in IIM phenotypes^{53,54}. Such studies have investigated possible disease associations of selected IIM phenotypes with many viral, bacterial, and parasitic infections⁵⁵; foods and dietary supplements⁵⁶; collagen and silicone implants^{57,58}; dozens of biologic agents and chemicals prescribed as drugs⁴⁶; seasonal variations¹⁰; birth date associations⁴²; exposure to ultraviolet light^{41,59,60}; and occupational exposures to dust, gases, or fumes^{61,62}.

A number of specific infectious agents are implicated in IIM pathogenesis on the basis of reported occurrences of suspected infection-induced disease or biologic plausibility from animal models⁵². Examples include hepatitis B virus in polymyositis⁶³ and dermatomyositis⁶⁴; hepatitis C virus in IBM⁶⁵; retroviruses, particularly human immunodeficiency virus (HIV) and human T-lymphotropic virus-1 (HTLV-1) in polymyositis⁶⁶, dermatomyositis⁶⁷, and IBM^{68,69}; *Toxoplasma* spp. and *Borrelia* spp. in polymyositis and dermatomyositis⁷⁰; and influenza, picornaviruses, and echovirus in polymyositis, dermatomyositis, and juvenile dermatomyositis⁵⁵. Case reports have documented associations between IIM development and medications, including D-penicillamine in polymyositis and dermatomyositis⁷¹; therapeutic cytokines, especially interferons⁷² and anti-TNF agents in dermatomyositis⁷³; and statins in polymyositis, dermatomyositis, necrotizing myopathy, and IIM with anti-HMGCR autoantibodies^{74–76}. On the basis of case series and animal models, vaccines (especially those containing aluminum hydroxide) have been hypothesized to be triggers of polymyositis, dermatomyositis, and focal forms of myositis that affect only a selected extremity or part of the body⁷⁷.

Because of the limitations of animal models and case reports, and despite the rarity and heterogeneity of the IIM, investigators have also assessed environmental risk factors by using epidemiologic approaches. The environmental associations already discussed have in some cases, but not in others, been supported by the few epidemiologic studies reported to date, which often addressed only a single exposure and have in many cases not been replicated. Those studies have sometimes had conflicting results, probably due to differing approaches and study populations. The epidemiologic studies to date reveal a number of preliminary environmental associations (Figure 2): the risk of IIM appears to be increased

after any infection and after gastrointestinal or respiratory tract infections or lung inflammation⁷⁸; the risk of polymyositis and dermatomyositis is decreased after upper respiratory tract infections⁷⁹; the risk of polymyositis or dermatomyositis is increased after excess physical exertion⁷⁹; there is no association with vaccines in polymyositis or dermatomyositis⁷⁹; smoking is a risk factor for anti-Jo-1 autoantibody-positive IIM (potentially interacting with *HLA-DRB1*03* to increase this risk, an effect similar to the HLA–smoking interaction seen in rheumatoid arthritis) and for myositis overlap syndromes⁸⁰; bovine collagen implants are associated with dermatomyositis⁵⁷; and group A Streptococcus infections are associated with juvenile dermatomyositis⁸¹. Perinatal factors in mothers, including air pollution, smoking, and occupational exposure to dust and/or solvent, have also been proposed as important risk factors for juvenile dermatomyositis in a small study⁸².

The compelling data that reveal a link between the environment and autoimmunity, as well as the remarkable increases in the incidence and prevalence of many autoimmune diseases for unknown reasons^{37,83–85}, underscore a critical need for both exploratory and confirmatory environmental investigations in this understudied field. Identifying factors that protect against disease is also important to decrease the prevalence of autoimmune disease. Much more work is needed in these areas. Enumerating genetic and environmental risk and protective factors in carefully defined disease phenotypes is an important first step for discovering gene–environment interactions that could lead to preventive strategies^{34,46}.

Immune-mediated disease mechanisms

Preliminary data suggest that many adaptive and innate immune mechanisms, as well as non-immune mechanisms, are involved in the development of the IIMs (Figure 3).

T cells in IIM

Several studies have demonstrated the presence of T cells and B cells, especially invasive, granzyme B and perforin-expressing cytotoxic CD8⁺ effector T cells, in the muscles of patients with polymyositis or IBM^{86,87}. The invasion of CD8⁺ effector cells into non-necrotic muscle fibres is considered a characteristic histological feature of polymyositis and IBM. The number of invading CD8⁺ and CD57⁺ T cells in the muscle correlates with the size of large granular lymphocyte populations in the blood in patients with IBM, and the autoimmune T cell expansion is proposed to evolve into a neoplastic-like or overtly neoplastic disorder of T cell aggressiveness⁸⁸.

In the 1990s, investigations demonstrated the presence of clonal $\gamma\delta$ T cells in the muscle of patients with polymyositis, and in 2012, investigators showed that these $\gamma\delta$ T cells recognize aminoacyl-tRNA synthetases (an MSA target), suggesting a potential link between $\gamma\delta$ T cells and autoantibody responses in autoimmune myositis^{89–91}. Patients with $\gamma\delta$ T cells in their muscle tissue seem to be more responsive to steroids compared to those without these cells in muscle^{89,92}. Muscles affected by myositis also contain some unique subsets of highly cytotoxic, apoptosis-resistant, pro-inflammatory T cells, such as CD28^{null} T cells (CD4+CD28^{null} and CD8+CD28^{null})^{93–95}. Persistence of CD244+ (CD28^{null}) T cells in

muscle tissue following immunosuppressive therapy is associated with poor outcome in patients with IIM^{95} .

The role of regulatory T (T_{reg}) cells (defined by FoxP3 expression) and the effect of these cells on inflamed muscle were investigated in adult and juvenile forms of myositis^{96,97}. It was proposed that FoxP3 T_{reg} cells counterbalance muscle inflammation in myositis. The presence of significant inflammation in the muscle of juvenile dermatomyositis patients despite a high proportion of T_{reg} cells in the milieu suggests that T_{reg} cell function is impaired. In a muscular dystrophy model, T_{reg} cells derived from the muscle express growth factors, such as amphiregulin, that act directly on muscle satellite cells in vitro and improve muscle repair in vivo⁹⁸.

B cells in IIM

Despite initial reports of the predominance of B cells in the perivascular regions on muscle biopsy in dermatomyositis⁷⁶, subsequent studies have demonstrated the presence of B cells, plasma cells, and immunoglobulin transcripts in the muscle of patients with polymyositis or IBM, indicating a humoral component in those disorders^{99–101}. Autoantibodies are present in more than half of all patients with IIM. The presence of MSAs, such as anti-Jo1, anti-Mi2, anti-cytosolic 5'-nucleotidase 1A (cN1A), or anti-HMGCR autoantibodies, are well described in patients with polymyositis, dermatomyositis, IBM, or necrotizing myopathy, respectively¹⁰². In addition to the prior work confirming many genetic and clinical associations with the classic myositis autoantibodies, studies have highlighted unique associations of newly identified autoantibodies with additional clinical phenotypes. For example, autoantibodies targeting melanoma differentiation-associated protein 5 (MDA5) are associated with mucocutaneous lesions and severe lung disease in patients with dermatomyositis; autoantibodies that bind to nuclear matrix protein 2 (NXP2) are associated with joint contractures and calcinosis in juvenile dermatomyositis; and autoantibodies to SAE, transcription intermediary factor 1γ (TIF1 γ), and transcription intermediary factor 1α (TIF1a) are associated with malignancy in dermatomyositis¹⁰³.

Although the pathological roles of autoantibodies in IIM are unclear, their associations with distinct genotypes, clinical phenotypes, and interferon patterns are strong. For example, the spectrum of clinical phenotypes in antisynthetase syndrome, a group of IIM defined by the presence of autoantibodies to aminoacyl-tRNA-synthetases that are involved in protein synthesis, includes interstitial lung disease, arthritis, Raynaud phenomenon, and mechanic's hands. Levels of anti-Jo1 autoantibodies are also strongly associated with clinical outcomes, suggesting that anti-Jo1 autoantibody levels might be a good biomarker for disease activity¹⁰⁴. Furthermore, animal and in vitro studies demonstrate that the Jo1 autoantigen, histidyl-tRNA synthetase, might also play a role in inducing myositis¹⁰⁵ or function as a chemokine¹⁰⁶.

The role of autoantibodies in causing muscle damage and dysfunction is controversial because most of the autoantigens are intracellular and thus not easily accessible to circulating autoantibodies; however, additional work focusing on their role is warranted given their importance as biomarkers of unique genetic, clinical, therapeutic, and outcome phenotypes. Although cause-and-effect relationships between autoantibodies and disease

phenotype and activity currently are not clear, the strong autoantibody associations with clinical phenotypes and disease severity are immensely useful for the diagnosis and prognosis of IIM.

Innate immune cells

Antigen processing and presentation by professional antigen-presenting cells, such as dendritic cells, are critical to initiating the adaptive immune response, and the muscle microenvironment of IIM is enriched with both myeloid and plasmacytoid dendritic cells^{107–111}. The relative proportion of lymphocytes and macrophages in skeletal muscle seems to vary in different clinical entities; for example, in anti-HMGCR myopathy, M2 (alternatively activated) macrophages predominate over CD4⁺ and CD8⁺ T cells and CD123⁺ plasmacytoid dendritic cells¹¹². A predominance of M2 macrophages in skeletal muscle is also seen in forms of inflammatory myopathy with abundant macrophages¹¹³. The presence of M2 macrophages is consistent with the known role of macrophages in skeletal muscle injury and repair¹¹⁴: M1 (classically activated) macrophages arrive later to sustain tissue healing¹¹⁵.

Inflammatory mediators

The high serum levels of creatine kinase and other molecules that are released from skeletal muscle cells in patients with IIM could represent danger-associated molecular patterns, which sometimes serve as endogenous Toll-like receptor ligands. Skeletal muscle, as well as muscle-infiltrating cells, express abundant innate immune receptors, including Toll-like receptors^{116–118}. Activation of innate immune receptors can lead to activation of NF- κ B signalling and pro-inflammatory cytokine and chemokine secretion, which, in turn, further recruit immune cells into a milieu that is already ripe for antigen processing and presentation by dendritic cells. These immune cells and cytokines can then further activate T helper 1 (T_H1), T helper 17 (T_H17), and T helper 1 (T_H2) cells, as well as CD8⁺ cytotoxic T cells and CD28^{null} cells, which potentially damage muscle cells. Cytokines produced by these T helper cell subsets induce macrophages to polarize into pro-inflammatory M1 or pro-resolution M2 phenotypes.

Both pro-inflammatory and anti-inflammatory cytokines, as well as CXC-chemokines and CC-chemokines, are expressed in IIM muscle; such mediators include T_{H1} cytokines (TNF, IFN γ , IL-12, and IL-2), T_{H2} cytokines (IL-4 and IL-13), T_{H1} 7 cytokines (IL-17, IL-22, IL-23, TNF-related weak inducer of apoptosis (TWEAK), and IL-6), T_{reg} cytokines (IL-10, transforming growth factor- β (TGF β)), and innate immune cytokines (IL-1 α , IL- β , and type I interferons (IFN α and IFN β)). These cytokines coordinate various innate and adaptive immune response pathways, and some of them have the potential to cause muscle damage and weakness depending on the stage of the disease^{119,120}. Similarly, both CXC-chemokines (CXC-chemokine ligand 9 (CXCL9), CXCL10) and CC-chemokines (CC-chemokine 2 (CCL2), CCL3, CCL4, CCL19, and CCL21) have a role in sustaining inflammatory responses in IIM muscle¹²¹.

In dermatomyositis, evidence from the past decade has demonstrated high levels of type 1 interferon in the muscles of patients, which is associated with perifascicular atrophy^{122–125}. Genes induced by type I interferons (either IFNa or IFN β) were overexpressed in the muscle, skin, and blood, of patients with dermatomyositis, and in some^{126–128}, but not all¹²⁹, of those studies the levels of these genes correlated with disease activity. Evidence suggests that myeloid dendritic cells are a major source of type 1 interferons in the muscle of dermatomyositis patients¹⁰⁸. Type I interferons affect immune cells either directly, through type I interferon receptor signalling, or indirectly, by inducing the production of chemokines, by inducing the secretion of cytokines such as IL-15 (which regulate natural killer (NK) cell and memory CD8⁺ T cell proliferation), by stimulating dendritic cells (which, in turn, activate naive T cells), or by inducing the differentiation of monocytes–macrophage lineage cells¹³⁰.

IFNβ can induce reactive oxygen species and mitochondrial damage in dermatomyositis¹³¹, which provides an important link to the cause of the functional impairment in this disorder. Induction of tissue inflammation and autoimmunity by IFNα involves direct toxic effects on tissue as well as provocation of destructive bystander immune responses¹³². IFNα mediates a long-lasting and preferential MHC class I overexpression in non-immune cells, such as human pancreatic beta cells and thyroid follicular cells, which usually lack MHC class I expression. This finding suggests that IFNα might amplify antigen presentation in type 1 diabetes and Hashimoto thyroiditis^{133,134}. The increased expression of MHC class I molecules on skeletal muscle cells of patients with IIM probably leads to increased susceptibility of the cells to cytotoxic T cell attack and endoplasmic reticular (ER) stress–mediated cell death^{135,136}. On the basis on these findings, neutralization of type 1 IFN has been explored as a treatment option in dermatomyositis and polymyositis¹³⁷.

Non-immune-mediated disease mechanisms

Apart from the inflammatory pathomechanisms discussed in the previous section, mounting evidence suggests that several non-immune-mediated mechanisms also operate in IIM (an overview of the topic has been provided elsewhere^{138,139}). In general, these mechanisms fuel inflammation via a positive feedback loop, affect muscle contraction and cause muscle weakness, imbalance muscular protein homeostasis, and lead to atrophy and mostly irreversible structural damage of muscle fibres. The suggestion that non-immune-mediated mechanisms are clinically relevant in IIM derives from a number of key findings: the muscular inflammation identified by muscle biopsy and MRI does not always correlate with the clinical severity; the effects of immunosuppressive treatments can be limited; and several non-inflammatory mechanisms, including cell stress and degenerative mechanisms are discussed in this section.

ER stress

ER stress is one of the best-studied elements of non-immune-mediated damage to skeletal muscle in all forms of IIM (overview in¹⁴⁰). ER stress mechanisms include the unfolded protein response (UPR) and the ER overload response (EOR). UPR is characterised by

upregulation of cyclic AMP-dependent transcription factor 6α (ATF6α), eukaryotic translation initiation factor 2a-kinase 3 (EIF2a kinase, also known as PERK), serine/ threonine-protein kinase/endoribonuclease IRE1 (IRE1a), and the ER chaperones endoplasmin (also known as GRP94) and 78-kDa glucose-regulated protein (GRP78, also known as BiP). The collective function of these molecules is to reduce the protein overload and subsequent accumulation of unfolded proteins in the ER. The second ER stress pathway, EOR, modulates inflammation by upregulating NF- κ B signalling. Both ER stress pathways are activated in the muscle in all forms of IIM, including IBM^{135,141,142}. Data from the past years suggest that ER stress might even contribute directly to muscular weakness in IIM (overview in¹⁴³). The NF- κ B pathway has been shown to be activated in IIM^{144,145}. At the same time, relevant molecules of the immunoproteasome such as β 1i and β 5i were present in the muscle of patients with IIM¹⁴⁶. The NOD-, LRR- and pyrin domain containing protein 3 (NLRP3) inflammasome has also been shown to be upregulated in dermatomyositis and polymyositis, and this was associated with elevated levels of IL-1β and IL-18¹⁴⁷. Since it is known that ER stress can induce the NLRP3 inflammasome in other cell systems¹⁴⁸, it is possible that ER stress is a crucial factor of muscle pathology in IIM by inducing molecules of the inflammasome and immunoproteasome pathways.

Other key factors

Free radicals are key factors in muscle fibre damage in all forms of $IIM^{149,150}$, and these molecules are speculated to contribute directly to muscle weakness¹⁴³. In a mouse model of chronic inflammation (mice overexpressing TGF β), muscular atrophy was mediated by inflammation, production of reactive oxygen species, mitochondrial damage, and caspase activation. The mitochondrial damage and muscle atrophy were efficiently downmodulated following red grape polyphenols supplementation despite continuous muscle inflammation¹⁵¹.

TNF-related apoptosis-inducing ligand (TRAIL) expression is upregulated and associated with autophagy and cell death in the skeletal muscle of patients with IIM¹⁵². In IIM muscle fibres, the expression of heat-shock proteins HSP70 and HSP90¹⁵³ and the alarmin high mobility group box protein 1 (HMGB1)¹⁵⁴ is increased compared with healthy controls; HMGB1 functions via Toll-like receptor-4 and is thought to mediate muscular inflammation and weakness^{155–157}. Consistent findings relating to increased HMGB1 expression have been reported in experimental models of autoimmune myositis in rodents¹⁵⁸. Dysregulation of HMGB1 might also be relevant for the regenerative potential of skeletal muscle in IIM, because HMGB1 is an important factor during recovery and is required for the patient to regain muscle function after severe damage¹⁵⁹ and, thus, could serve as a prognostic marker in severe IIM¹⁶⁰.

Inclusion body myositis

The largest body of evidence for non-immune-mediated mechanisms in IIM is available for IBM¹⁶¹. The pathogenesis of IBM includes many pathways involved in protein homeostasis and cell stress mechanisms, and several of these pathways seem to be linked directly to inflammation.

Protein homeostasis and the heat shock response—A variety of unwanted and defective proteins that should be removed from the cell, including β -amyloid and its associated proteins, can accumulate in the muscle of patients with IBM^{162–164}. In IBM, intracellular accumulation of abnormal or no-longer-needed proteins in muscle fibres is hypothesized to cause or aggravate cell stress pathways, thus leading to structural damage and weakness of the fibres. In patients with IBM, proteomic analysis of vacuolated fibres identified FYVE domains and coiled-coil domain–containing protein 1 (FYCO1) as a novel component in the rimmed vacuoles²⁷. This finding was associated with a missense variant of the *FYCO1* gene in 11% of IBM patients (see above), which supports the hypothesis of impaired autophagic activity in IBM (see below for details).

HSP expression has been demonstrated in the muscles of patients with IBM and in human muscle fibres cultured in experimental conditions that mimic either inflammatory or degenerative aspects of IBM pathology^{165,166}. One of the most commonly expressed HSPs in IBM, crystallin α -B chain (also known as α B-crystallin or CRYAB or HSPB5), seems to be an early element in the pathological cascade, as this protein is upregulated in healthy muscle fibres^{165,167}. Heat-shock factor protein 1 (HSF1) can ameliorate cell stress caused by aggregation of TAR DNA binding protein 43 (TDP43) in muscle cells¹⁶⁸. In cell culture and animal models with pathologic features of IBM, treatment with arimoclomol, a modulator of the heat-shock response, protected skeletal muscle cells against protein accumulation and cell stress¹⁶⁹. The same study showed that arimoclomol was safe and well-tolerated in patients with IBM. Collectively, these data suggest that protein dyshomeostasis is an important non-immune element in the pathology of IBM and that this process is associated with a heat-shock response that could be a suitable target for future clinical trials.

Dysregulation of autophagy—Several lines of evidence have demonstrated malfunction of the autophagic machinery in IBM^{170,171}. Macroautophagy is active during the accumulation of β -amyloid in vacuoles (although it is not known yet whether macroautophagy causes β -amyloid to accumulate)¹⁷², and this process depends on extracellular signal-regulated kinase (ERK) signalling¹⁷³. Other autophagy adaptor molecules have been implicated in IBM, including sequestosome 1 (SQSTM1; also known as p62)¹⁷⁴, NBR1^{175,176}, and nuclear factor erythroid 2-related factor 2 (NRF2; also known as NFE2L2)¹⁷⁷. Collectively, these data indicate that autophagy is a relevant mechanism in IBM pathology and provide the rationale for a recently completed, placebo-controlled clinical trial of rapamycin, an immunosuppressant that activates macroautophagic activity, in patients with IBM (results not yet published)¹⁷⁸.

Mitochondrial abnormalities and free radicals—Apart from inflammation and protein accumulation, mitochondrial abnormalities, such as cyclo-oxygenase-deficient muscle fibres, are hallmarks of IBM. Several mitochondrial defects have been demonstrated in the muscle of patients with IBM¹⁷⁹. Such mitochondrial changes are associated with oxidative damage¹⁸⁰, inflammatory mediators¹⁸¹, and functional impairment of muscle strength¹⁸². Signs of mitochondrial dysfunction have also been shown in a mouse model of IBM, in which mice overexpressing human amyloid precursor protein develop IBM-like pathological features owing to the accumulation of amyloid precursor protein in the skeletal

muscle¹⁸³ and in human muscle cells following adenovirus-mediated upregulation of amyloid precursor protein in vitro.

In addition to the reports of free radical generation in IIM, production of nitric oxide has been demonstrated in IBM in association with accumulation of β -amyloid and inflammation in skeletal muscle as well as in different cell culture systems that mimic relevant aspects of IBM^{184,185}.

Targeting non-immune-mediated pathways

Non-immune-mediated mechanisms are relevant in all forms of IIM and can lead to structural damage of muscle fibres, cause direct weakness of the muscle, or reciprocally fuel inflammatory cell stress pathways. Many of these pathways are far from being understood and present possibly novel areas of mechanistic studies and therapeutic approaches. One of the novel treatment strategies in IBM is to regulate the function of protective HSPs. Future treatment directions should include therapies that scavenge free radicals and/or target other damage signals, such as alarmins and non-inflammatory mediators. Such therapies could reduce structural damage to muscle fibres and ameliorate weakness, particularly when these features are mediated by a functional impairment of muscle homeostasis and energy metabolism, such as mitochondrial dysfunction.

Conclusions

Studies in the past decade have elucidated possible genetic and environmental risk factors, as well as possible immune and non-immune mechanisms, that result in the development of IIM phenotypes. Yet, current paradigms are often biased by certain assumptions and use non-standardised disease or phenotype definitions that can limit our understanding and result in the assessment of different entities by different investigators. Current studies have emphasised the importance of using mutually exclusive and stable phenotypes to minimize confounding factors and to allow for greater power by using more homogeneous groups and smaller sample sizes, as are needed for rare diseases such as the IIMs. Multidisciplinary IIM collaborative study groups have played a key role in developing consensus on how to define and study IIM phenotypes (see the Reviews by Lundberg et al. *<citation>* and Rider et al. *<citation>* on classification and outcome assessment, respectively, in this Issue).

In the future, emphasis needs to be placed on multidisciplinary collaborative investigations of genetic and environmental risk factors and their interactions, as well as pathogenic mechanisms in homogeneous, well-defined phenotypes, utilizing the many IIM registries and repositories that have been developed to allow for the most cost-effective strategies¹⁸⁶. More investment in these areas seems appropriate to develop preventative strategies and to allow for innovative approaches to treatment as new pathways to disease are discovered.

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Glossary terms

Rimmed vacuoles

a space within the cytoplasm of a muscle cell with a purplish staining rim on trichrome staining

Functional annotation

characterizing the function assigned to each gene product or genetic variant

Expression quantitative trait locus (eQTL)

a genomic locus that regulates gene expression

Macroautophagy

a process in which cellular contents are degraded by lysosomes or vacuoles and recycled

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KEY POINTS

- Multiple independent associations within the HLA 8.1 ancestral haplotype are the strongest genetic risk factors for IIMs (idiopathic inflammatory myopathies).
- Non-HLA associations overlap with those for other autoimmune diseases, whereas some genetic risk factors are unique to IIM phenotypes, suggesting that these phenotypes have different pathophysiologies.
- In addition to drug-induced myositis, epidemiologic data support a role for infections, preceding lung disease, physical exertion, collagen implants, ultraviolet radiation, and smoking in the development of IIM phenotypes.
- Although the disease mechanisms for IIM phenotypes are ill defined, the innate (including cytokines and chemokines) and adaptive immune systems (including autoantibodies, and antigen-specific T cells) are likely involved.
- Several non-immune-mediated mechanisms contribute to IIM pathogenesis, including cell stress pathways, free radicals, altered energy metabolism, protein homeostasis, and mitochondrial damage.
- Multidisciplinary collaborative approaches, focused resources, and better investigative tools are needed to define additional risk and protective factors and pathogenic mechanisms, to cure and prevent the development of IIM phenotypes.

Box 1:

The Idiopathic Inflammatory Myopathies

- The idiopathic inflammatory myopathies (IIM), or myositis, are a heterogeneous group of rare systemic disorders that might involve multiple organ systems but are defined by chronic muscle inflammation.
- The major IIM phenotypes are dermatomyositis, polymyositis, and necrotizing myopathies, which have a female predominance¹⁸⁷, and inclusion body myositis, which has a male predominance¹⁸⁸.
- Childhood-onset and adult-onset forms of IIMs share many clinical, pathologic, and genetic features but differ in the frequency of specific phenotypes and in response to therapies and prognosis.
- Certain autoantibodies directed against translational and transcriptional components are found only in the IIMs and define unique clinical, genetic, and prognostic groups.
- Both immune and non-immune pathways contribute to muscle damage and weakness.
- Therapy in IIMs includes immunosuppressive agents to decrease the inflammation and rehabilitation with exercise to strengthen remaining muscle.

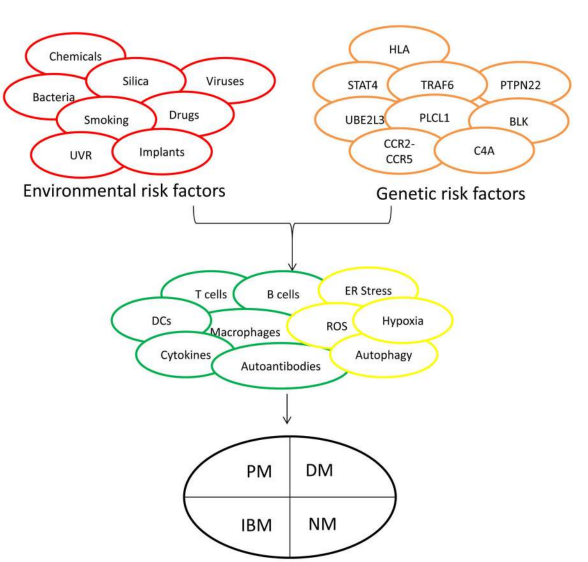


Figure 1. Possible pathways to idiopathic inflammatory myopathy phenotypes

The idiopathic inflammatory myopathies (IIM) probably consist of multiple phenotypes, each of which might be defined by unique combinations of symptoms, signs, and laboratory abnormalities. The major IIM phenotypes—polymyositis (PM), dermatomyositis (DM), necrotizing myopathy (NM), and inclusion body myositis (IBM)—are shown in the black circle at the bottom. Each phenotype could result from different pathogenic mechanisms, as represented by green circles (showing immunologic processes) and yellow circles (showing non-immunologic processes) in the center of the figure, because of the interactions between genetic risk factors (orange circles) and environmental risk factors (red circles). Some combinations of genotypes and environmental exposures induce certain mechanisms and disease phenotypes, whereas other combinations might have no effect or could be protective. C4A, complement 4A; DCs, dendritic cells, ER, endoplasmic reticulum; HLA, human leukocyte antigen; ROS, reactive oxygen species; UVR, ultraviolet radiation.

Exposure	IIM phenotype	Comments	OR or RR (95% CI)	OR or RR (95% CI)	Re
Infectious agents					
Infections more than 1 year prior to IIM diagnosis All	IIM	Swedish nationwide registry	-	1.5 (1.2 - 1.9)	14
GI	IIM			1.9 (1.1 - 3.5)	14
Respiratory	IIM			2.3 (1.8 - 3.3)	14
URI one year prior to onset	PM, DM	Questionnaire, case-sibling control		0.35 (0.17 - 0.71)	14
Group A Streptococcus	JPM, JDM	Case-control household exposures		- 2.73 (1.14 - 6.53)	14
Parvovirus	JDM	No association seen -		0.35 (0.14 - 0.90)	14
Non-infectious agents					
Physical exertion	PM, DM	Questionnaire, case-sibling control		3.9 (1.8 - 8.2)	14
Vaccines	PM, DM	Questionnaire, case-sibling control	_	1.0 (0.5 - 2.2)	14
Bovine collagen implants	DM	Cohort study		5.05 (2.31 - 9.59)	15
Smoking	Anti-Jo-1 autoantibodies	Strongest association seen in Hungarian IIM patients, and interaction with HLA DR3		3.94 (1.53 - 9.89)	2
Smoking	CTD-overlap myositis	European cohort comparing clinical phenotypes	· · · · · · · · · · · · · · · · · · ·	1.44 (1.08 - 1.90)	15
		0.1	1	10	
Inflammatory lung disease at study inclusion	IIM	Swedish inpatient registry		5.3 (1.3-21.7)	14
Maternal smoking during pregnancy	JDM	Brazilian case-control study		13.26 (1.21 – 144.29)	15
Maternal occupational exposure to chalk dust or gasoline vapor	JDM	Brazilian case-control study		35.39 (1.97 - 632.80)	15
Maternal exposure to atmospheric carbon monoxide	JDM	Brazilian case-control study		12.21 (1.28 - 115.96)	15

Figure 2. Epidemiologic investigations of environmental agents and IIMs

Infectious and non-infectious agents associated with IIM phenotypes from epidemiologic studies suggest a wide array of risk factors with different strengths of association. 95% CI, 95% confidence interval; CTD, connective tissue disease; DM, dermatomyositis; GI, gastrointestinal; HLA, human leukocyte antigen; IIM, idiopathic inflammatory myopathies; JDM, juvenile dermatomyositis; JPM, juvenile polymyositis; OR, odds ratio; PM, polymyositis; RR, relative risk; URI, upper respiratory infection.

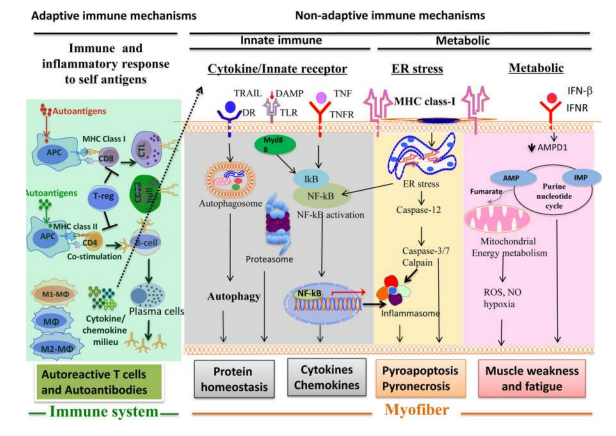


Figure 3.

Infectious and non-infectious agents evaluated in epidemiological studies suggest a wide array of risk and protective factors with different strengths of association with idiopathic inflammatory myopathy (IIM) phenotypes. CTD, connective tissue disease; OR, odds ratio; RR, relative risk. **a.** Infection in the year before IIM diagnosis. **b.** Infection in the year before IIM onset. **c.** Inflammatory lung disease at study inclusion.

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IIM phenotype	Gene or allele (associated SNP or amino acid position)	p value	OR (95% CI)	Study type	Putative function	Associations with other autoimmune disease	Reference
IIM	HLA -DRB1*03:01 †	2.58×10 ⁻¹³⁵	1.88 (1.68–2.11)	Immunochip & HLA imputation	Regulation of immune system	Type I diabetes, SLE, Crohn's disease	13
	HLA -B*08:01 †	3.23×10^{-14}	1.58 (1.41–1.78)	Immunochip & HLA imputation	Regulation of immune system	Unknown	13
	<i>STAT4</i> (rs4853540)	1.57×10 ⁻⁶	0.83 (0.77–0.89)	Immunochip	T cell differentiation	Celiac disease, Crohn's disease, IBD	13
	TRAF6 (rs570676)	9.42×10 ⁻⁶	0.87 (0.82–0.92)	Immunochip	TNF receptor signalling	RA	13
	UBE2L3 (rs5754467)	4.67×10^{-7}	1.21 (1.12–1.30)	Immunochip	NF- kB signalling	Celiac disease, psoriasis SLE	13
IIM (Japanese population)	HLA-DRB1*08:03	0.02	1.9 (1.1–3.2)	Candidate gene	Regulation of immune system	Allergic rhinitis	15
Polymyositis	HLA-DRB1 $*03:01$ ^{\dagger}	6.11×10^{-80}	1.99 (1.67–2.36)	Immunochip & HLA imputation	Regulation of immune system	Type I diabetes, SLE, Crohn's disease	13
	HLA-B $*08:01^{\dagger}$	4.17×10 ⁻⁹	1.71 (1.43–2.05)	Immunochip & HLA imputation	Regulation of immune system	Unknown	13
	<i>HLA-DRB1</i> (asparagine 77)	1.65×10 ⁻⁸⁰	2.93 (2.53–3.17)	Immunochip & HLA imputation	Regulation of immune system	Type I diabetes, autoimmune thyroid disease	13
	PTPN22 (rs2476601)	7.90×10^{-11}	1.58 (1.38–1.81)	Immunochip	T cell signalling	RA, SLE, type I diabetes, autoimmune thyroid disease	13
Adult dermatomyositis	HLA-B*08:01	2.46×10 ⁻⁴²	1.90 (1.66–2.17)	Immunochip & HLA imputation	Regulation of immune system	Unknown	13
	<i>HLA-DQB1</i> (alanine 57)	1.29×10^{-12}	1.62 (1.44–1.83)	Immunochip & HLA imputation	Regulation of immune system	Type I diabetes	13
Adult dermatomyositis	HLA-DQA1*01:04	0.01	2.58 (1.18–5.64)	Candidate gene	Regulation of immune system	Unknown	16
(Cumos population)	HLA-DRB1*07	0.01	2.26 (1.12-4.59)	Candidate gene	Regulation of immune system	Unknown	16
Juvenile dermatomyositis	HLA-DRB1*03:01	7.91×10- ¹⁴	1.90 (1.61–2.22)	Immunochip & HLA imputation	Regulation of immune system	Type I diabetes, SLE, Crohn's disease	13
	C4A deficiency	8.2×10 ⁻⁶	3.00 (1.87–4.79)	Candidate gene	Classical complement pathway	SLE, Bechet syndrome	24
Juvenile and adult dermatomyositis	PLCL1 (rs7572733)	6.18×10 ⁻⁶	0.80 (0.72–0.88)	GWAS & candidate gene	Inositol phospholipid signalling	SLE	10

BLK (xc23:64.0)BCK (xc23:64.	IIM phenotype	Gene or allele (associated SNP or amino acid position)	p value	OR (95% CI)	Study type	Putative function	Associations with other autoimmune disease	Reference
$HLA-DRB/r030/t$ $S.77k10^{34}$ $T/73(5.8-10.95)$ Immunchip & HLARegulation of immue systemType I diabetes. SLE, Tohn's disease $DRB/r0.0/t$ $1.56k10^{-6}$ $1.56k10^{-6}$ $464(3.3.4.6)$ Immunchip & HLARegulation of immue systemPions' disease $DRB/r13.0/t$ $3.28k10^{-6}$ $3.9(2.14-1.72)$ Immunchip & HLARegulation of immue systemPions' disease $DRB/r13.0/t$ $3.28k10^{-6}$ $3.9(2.14-1.72)$ Immunchip & HLARegulation of immue systemPions' disease $DRB/r13.0/t$ $3.28k10^{-6}$ $3.9(2.14-1.72)$ Immunchip & HLARegulation of immue systemPions' disease $DRB/r13.0/t$ $1.95k10^{-6}$ $3.8(2.80-5.29)$ Immunchip & HLARegulation of immue systemPions' disease $DRB/r13.0/t$ $1.95k10^{-6}$ $3.8(2.80-5.29)$ Immunchip & HLARegulation of immue systemPions' disease $DRB/r13.0/t$ $1.95k10^{-6}$ $0.42(0.29-0.60)$ Immunchip & HLARegulation of immue systemPions' disease $DRB/r13.0/t$ $N(A)$ $N(A)$ $D(A)$ $D(A)$ Regulation of immue systemPions' disease $DOP PioneN(A)N(A)D(A)D(A)D(A)D(A)D(A)D(A)DOP PioneN(A)N(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)N(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)D(A)$		BLK (rs2736340)	0.000065	1.25 (1.12–1.40)	GWAS	B cell signalling and development	RA, SLE	10
$BRJ^*\eta;\eta;\eta$ 1.56×10^{-6} $6.4(3.33-6.4)$ Imunochip & HLARegulation of immue systemR $DRBJ^*J;\eta;\eta$ $3.28\times10^{*6}$ $3.19(2.14-4.72)$ imputationRegulation of immue systemR $DRBJ^*J;\eta;\eta$ $3.28\times10^{*6}$ $3.19(2.14-4.72)$ imputationRegulation of immue systemR $DRBJ^*J;\eta;\eta$ $3.28\times10^{*6}$ $3.19(2.14-4.72)$ imputationRegulation of immue systemR $HLA-DRBJ(tyosine 26)$ 1.19×10^{-16} $3.83(2.80-5.29)$ immuochip & HLARegulation of immue systemR VCP^{\pm} 1.99×10^{-6} $3.83(2.80-5.29)$ immuochip & HLARegulation of immue systemR VCP^{\pm} NA NA NA NA Targeted sequencingProtein homeostasisIIA, ulcrative colits, VCP^{\pm} NA NA NA NA Targeted sequencing NF -R signalling ALS^{K} VCP^{\pm} NA NA NA Targeted sequencing NF -R signalling ALS^{K} VCP^{\pm} NA NA NA Targeted sequencing NF -R signalling ALS^{K} VCP^{\pm} NA NA NA NA NA NA NA VCP^{\pm} NA NA NA NA NA NA VCP^{\pm} NA NA NA NA NA NA VA NA NA NA NA NA NA VCA NA NA NA NA NA NA NA NA <td< td=""><td>IBM</td><td>HLA-DRB1*03:01†</td><td>5.77×10⁻³⁴</td><td>7.97 (5.88–10.95)</td><td>Immunochip & HLA imputation</td><td>Regulation of immune system</td><td>Type I diabetes, SLE, Crohn's disease</td><td>14</td></td<>	IBM	HLA -DRB1*03:01 †	5.77×10 ⁻³⁴	7.97 (5.88–10.95)	Immunochip & HLA imputation	Regulation of immune system	Type I diabetes, SLE, Crohn's disease	14
$BRI^*/3.01^{\dagger}$ $3.28\times10^{\circ}$ $3.19(2.14-17)$ Immuochip & HLARegulation of immue systemPrimary sclerosing $HLA-DRB1(tyrosine 26)$ 1.19×10^{-16} $3.83(2.80-5.29)$ Immuochip & HLARegulation of immue systemBIA $HLA-DRB1(tyrosine 26)$ 1.19×10^{-16} $3.83(2.80-5.29)$ Immuochip & HLARegulation of immue systemSLE $CCR2-CCR5 region$ $1.93\times10^{\circ6}$ $0.42(0.29-0.60)$ Immuochip & HLARegulation of immue systemSLE VCp^{\ddagger} NA NA NA $Targeted sequencingProtein homeostasisIBMPFD^6A_{1S}SVCp^{\ddagger}NANANATargeted sequencingN-\kappa B signallingALS^{6}ALS^{6}VCp^{\ddagger}NANANANANANANAALS^{6}ALS^{6}ARS.1 alleles^{4}(HLA-0.003N/ANANANAAutohagic adaptor proteinUnhousARS.1 alleles^{4}(HLA-0.003N/ANANANANANANANAALS^{6}ARS.1 alleles^{4}(HLA-0.003N/ANA$		DRB1*01:01 [†]	1.56×10^{-16}	4.64 (3.33–6.49)	Immunochip & HLA imputation	Regulation of immune system	RA	14
$HLA-DRBI (tyrosine 26)$ 1.19×10^{-6} $3.3 (2.80-5.29)$ immunochip & HLARegulation of immune system SLE $CCR2-CCR5 region$ 1.93×10^{-6} $0.42 (0.29-0.60)$ immunochipBinds pro-inflammatory $IIA, ulcerative colits, china chinaVCp^{\ddagger}N/AN/AN/ATargeted sequencingProtein homeostasisIIA, ulcerative colits, chinaVCp^{\ddagger}N/AN/AN/ATargeted sequencingProtein homeostasisIIA, ulcerative colits, chinaVCp^{\ddagger}N/AN/AN/ATargeted sequencingN-K signallingA_LS^SFYCOI0.003N/AN/ATargeted sequencingN-K signallingA_LS^SAHB, 1alee^A/HLA-0.003N/ANole-exoneAutohagic adaptor proteinUnknownB_{S001, HLA-}0.003N/AN/ANele-exoneA_LS^SA_LS^SB_{R1}^{*}(32.0)und HLA B_{R2}^{*}(3.0)N/AN/ANele-exoneA_LS^SB_{R1}^{*}(32.0)und HLA B_{R2}^{*}(3.0)N/AN/ANele-exoneA_LS^SB_{R1}^{*}(32.0)und HLA B_{R2}^{*}(3.0)N/AN/AN/AN/AB_{R2}^{*}(3.0)ULA-DRBI^{*}(1.0)U(A, 0.3.1, 4)N/AN/AN/AN/AHLA-DRBI^{*}(1.0)U(A)U(A, 0.3.1, 4)U(A, 0.3.1, 4)N/AN/AN/AHLA-DRBI^{*}(1.0)U(A)U(A)U(A)U(A, 0.3.1, $		DRB1*13:01 [†]	3.28×10 ⁻⁸	3.19 (2.14-4.72)	Immunochip & HLA imputation	Regulation of immune system	Primary sclerosing cholangitis	14
$CR2-CCR5$ region 1.93×10^{-6} $0.42(0.29-0.60)$ ImmuocipBinds pro-inflammatoryIA, ulcrative colits, celia disease VCP^{\ddagger} N/A N/A N/A N/A N/A $Rageted sequencingProtein homeostasisIA_{\rm sefe} diseaseVCP^{\ddagger}N/AN/AN/ARageted sequencingProtein homeostasisIA_{\rm sefe}R_{\rm sefe}SQSTMI^{\ddagger}N/AN/ARageted sequencingNF+RB signallingA_{\rm LS}^{\$}A_{\rm LS}^{\$}FYCOI0.003N/AN/AWhole-exomeAutohagic adaptor proteinMenoneHR1 alleles^A(HLA 0.003N/AWhole-exomeAutohagic adaptor proteinMenoneB^{*}0R:01, HLA 0.003N/AWhole-exomeAutohagic adaptor proteinMenoneB^{*}0R:01, HLA 0.003N/AWhole-exomeAutohagic adaptor proteinMenoneB^{*}0R:01, HLA B^{*}0R:01, HLA B^{*}0R:01, HLA Regulation of immune systemIhe InhownB^{*}0R:01, HLA 1.20\times 10^{-83}N/AWhole-exomeRegulation of immune systemIhe InhownB^{*}0R:01, HLA 1.20\times 10^{-83}N/AGWAS \& HLARegulation of immune systemIhe InhownB^{*}0R:01, HLA 1.20\times 10^{-81}1.20\times 10^{-81}Ihe InhownIhe InhownIhe InhownB^{*}0R:01, HLA 1.20\times 10^{-81}Ihe InhownIhe InhownIhe InhownIhe InhownB^{*}0R:01, HLA Ihe Inhown$		HLA-DRB1 (tyrosine 26)	1.19×10^{-16}	3.83 (2.80–5.29)	Immunochip & HLA imputation	Regulation of immune system	SLE	14
VCP^{\ddagger} N/AN/ATargeted sequencingProtein homeostasisIBMPFD $^{\sharp}$ ALS $^{\$}$ VCP^{\ddagger} N/AN/ATargeted sequencingProtein homeostasisIBMPFD $^{\sharp}$ ALS $^{\$}$ $FYCO1$ N/AN/ATargeted sequencingNF-xB signalling $ALS^{\$}$ $FYCO1$ 0.003N/AWhole-exomeAutophagic adaptor proteinUnknown $AB8.1 alleles^A(HLA-)$ 0.003N/AWhole-exomeAutophagic adaptor proteinUnknown $B^{\$}08.01, HLA-$ U20x10- ⁸² N/AWhole-exomeRegulation of immune systemUnknown $DOB 1*02.01 and HLA-$ U20x10- ⁸² N/AGWAS & HLARegulation of immune systemType I diabetes, SLE, $DOB 1*02.01 and HLA U20-6$ 10.4(3.6-31.4)Candidate geneRegulation of immune systemType I diabetes, SLE, $HLA-DRB1*01.01$ 1.2×10^{-6} $10.4(3.6-31.4)$ Candidate geneRegulation of immune systemIA $HLA-DRB1*07.01$ 0.01 N/ACandidate geneRegulation of immune systemIA		CCR2-CCR5 region	1.93×10 ⁻⁶	0.42 (0.29–0.60)	Immunochip	Binds pro-inflammatory chemokines	JIA, ulcerative colitis, celiac disease	14
$SQSTMT^{\ddagger}$ N/AN/ATargeted sequencingNF-kB signalling $ALS^{\$}$ $FYCOI$ 0.003N/AWhole-exomeAutophagic adaptor proteinUnknown $H81.$ alleles^A(HLA- $B*08.01, HLA-$ 0.003N/AWhole-exomeAutophagic adaptor proteinUnknown $AH8.1$ alleles^A(HLA- $DOB1*02.01$ and HLA- $DOB1*02.01$ and HLA-1.20x10^{-82}N/AGWAS & HLARegulation of immune systemCinhor site ase $DOB1*02.01$ and HLA- $DOB1*02.01$ and HLA-1.20x10^{-82}N/AGWAS & HLARegulation of immune systemType I diabetes, SLE, Cinhor site ase $HLA-DRB1*11.01$ 1.2x10^{-6}10.4(3.6-31.4)Candidate geneRegulation of immune systemIA $HLA-DRB1*11.01$ 1.2x10^{-6}10.4(3.6-31.4)Candidate geneRegulation of immune systemIA $HLA-DRB1*07.01$ 0.01N/ACandidate geneRegulation of immune systemIA		VCP^{\ddagger}	N/A	N/A	Targeted sequencing	Protein homeostasis	IBMPFD [§] , ALS [§]	26,27
FYCO10.003N/AWhole-exomeAutophagic adaptor proteinUnknownAH8.1 alleles^A(HLA- B*08.01, HLA- DOB1*02.01 and HLA- 1.20×10^{-82} N/AGWAS & HLARegulation of immune systemType I diabetes, SLE, Crohn's disease $B*08.01, HLADOB1*02.01 and HLA-DRB1*03.011.20\times10^{-82}N/AGWAS & HLARegulation of immune systemType I diabetes, SLE,Crohn's diseaseHLA-DRB1*02.011.2\times10^{-6}10.4(3.6-31.4)Candidate geneRegulation of immune systemJIAHLA-DRB1*11.011.2\times10^{-6}10.4(3.6-31.4)Candidate geneRegulation of immune systemJIAHLA-DRB1*07.010.01N/ACandidate geneRegulation of immune systemJIA$		$_{\#}^{IWLSOS}$	N/A	N/A	Targeted sequencing	NF-xB signalling	ALS	26,27
AH8.1 alleles^{A}(HLA-) B*08.01, HLA- DOB1*02.01 and HLA- DCB1*02.01 and HLA-1.20×10 ⁻⁸² B*08.01, HLA- DCB1*02.01 and HLA- B*08.01)NAGWAS & HLA imputationRegulation of immune system Crohn's disease Crohn's disease HLA-DRB1*11:011.20×10 ⁻⁸² 1.2×10 ⁻⁶ NAGWAS & HLA imputationRegulation of immune system Begulation of immune systemType I diabetes, SLE, Crohn's diseaseHLA-DRB1*11:01 1.2×10^{-6} $10.4 (3.6-31.4)$ Candidate geneRegulation of immune systemJIAHLA-DRB1*07:01 0.01 N/ACandidate geneRegulation of immune systemParcreatitis		FYCOI	0.003	N/A	Whole-exome sequencing	Autophagic adaptor protein	Unknown	26,27
HLA - $DRB1*11:01$ 1.2×10^{-6} 10.4 ($3.6-31.4$)Candidate geneRegulation of immune system JIA HLA - $DRB1*07:01$ 0.01 N/A Candidate geneRegulation of immune systemPancreatitis	Anti-Jol positive IIM	AH8.1 alleles ^A (<i>HLA-</i> <i>B*08.01, HLA-</i> <i>DQB1*02.01 and HLA-</i> <i>DRB1*03.01</i>)	1.20×10 ⁻⁸²	N/A	GWAS & HLA imputation	Regulation of immune system	Type I diabetes, SLE, Crohn's disease	11
HLA-DRB1*07:01 0.01 N/A Candidate gene Regulation of immune system Pancreatitis	Anti-HMGCR positive adult myopathy	HLA-DRB1*11:01	1.2×10 ⁻⁶	10.4 (3.6–31.4)	Candidate gene	Regulation of immune system	JIA	19,20
	Anti-HMGCR positive juvenile myositis	HLA-DRB1*07:01	0.01	N/A	Candidate gene	Regulation of immune system	Pancreatitis	19,20

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95% CI, 95% confidence interval; AH8.1, 8.1 ancestral haplotype; ALS, amyotrophic lateral sclerosis; GWAS, genome-wide association study; HMGCR, anti-3-hydroxy-3-methylglutaryl-coenzyme A reductase; IBD, inflammatory bowel disease; IBM, inclusion body myositis; IBMPFD, IBM with Paget's disease of bone and frontotemporal dementia; IIM, idiopathic inflammatory myopathy; JIA, juvenile idiopathic arthritis; N/A, not applicable; NF-xB, nuclear factor xB; OR, odds ratio; RA, rheumatoid arthritis; SLE, systemic lupus erythematosus.

 $\dot{\tau}_{\rm Independent}$ effects, after conditioning.

 $\dot{\tau}^{t}$ Multiple rare variants reported.

 \S Not an autoimmune disease.

^{Λ} Considered together as a haplotype.