

Paths to understanding the genetic basis of autoimmune disease

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Some people inherit an unfortunate combination of genetic sequences, such that exposure to an external trigger causes their immune response to turn on their own tissues. Although mutations in a single gene can cause autoimmunity, most autoimmune diseases are associated with several sequence variants. Marked advances in genetic resources and tools are now making it possible to identify the sequence variants that contribute to autoimmune diseases — promising a better understanding of how we normally remain tolerant of our own tissue components, and how this goes wrong in autoimmune disease.

Autoimmune diseases are major causes of morbidity and mortality throughout the world. Many of these diseases tend to be difficult or impossible to cure, for the obvious reason that the focus of the immune response — self antigens — cannot be eliminated. The physical, psychological and economic burden of these diseases is especially devastating because they often attack young adults. The problem is also compounded by the failure of conventional cellular immunological analyses to shed much light on the pathogenic mechanisms. Recently developed therapies, such as tumour necrosis factor (TNF) antagonists, have had some remarkable successes, but these treatments target resulting organ damage and not the (usually unknown) underlying causes. The realization that the development of autoimmunity is strongly influenced by inherited polymorphisms (or DNA sequence variations) brings hope that understanding the genetics of autoimmune diseases will teach us about the causal derangements, and perhaps lead to new therapeutic strategies. Here, we summarize our current understanding of the genetic basis of autoimmunity. We emphasize principles, rather than attempt to list all the reported disease-associated polymorphisms. A brief introduction to the mechanisms of self tolerance and its breakdown provide the foundation for the subsequent discussion of the genetics of autoimmune diseases.

Self tolerance and autoimmunity

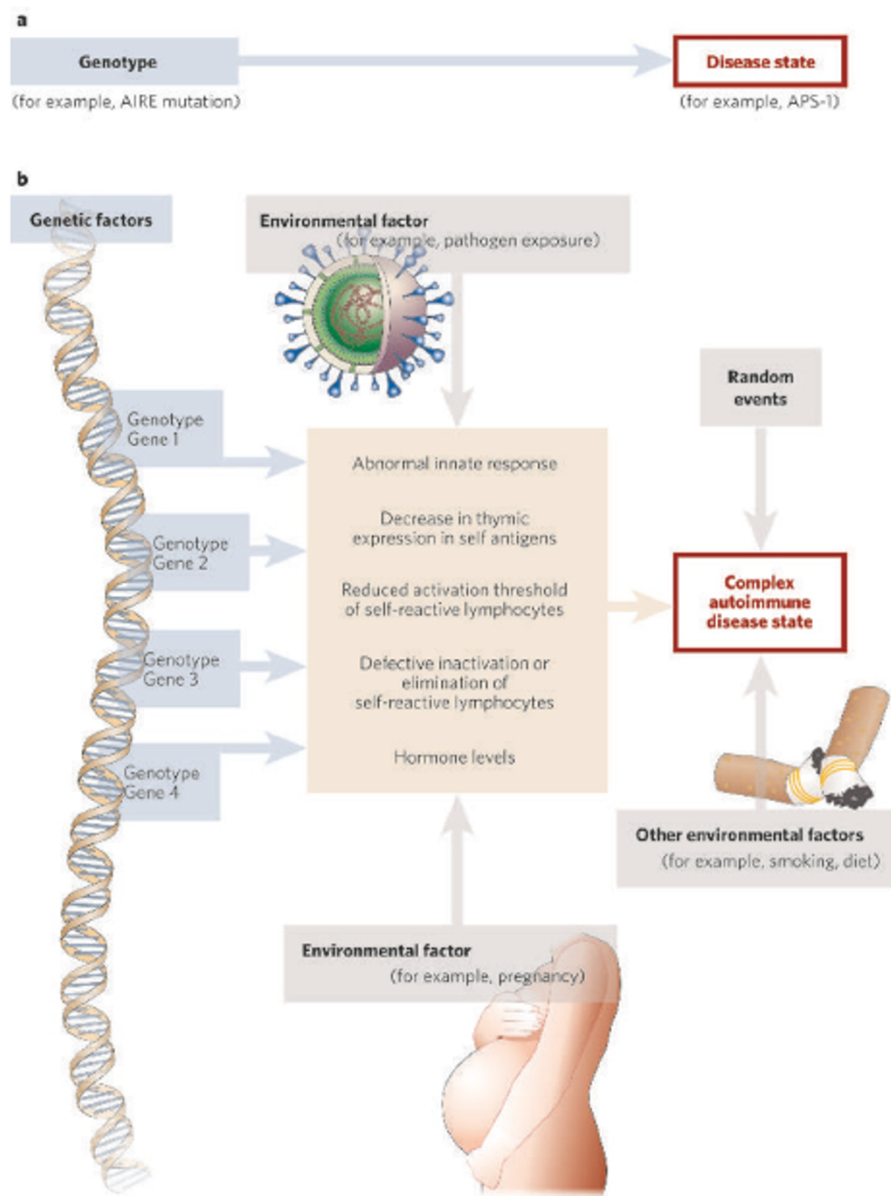
All individuals are tolerant of their own potentially antigenic substances, and failure of self tolerance is the fundamental cause of autoimmunity. The mechanisms of self tolerance have been worked out in considerable detail in animal models, and are best understood for CD4⁺ T cells. Self tolerance can be divided into central tolerance and peripheral tolerance. In central tolerance, immature lymphocytes that happen to recognize self antigens in generative lymphoid organs (the bone marrow for B cells and the thymus for T cells) die by apoptosis; in peripheral tolerance, mature self-reactive lymphocytes encounter self antigens in peripheral tissues and are killed or shut off. The principal mechanisms of peripheral tolerance are anergy (functional unresponsiveness), deletion (apoptotic cell death), and suppression by regulatory T cells¹. These mechanisms are described in more detail in the reviews in this issue by Goodnow *et al.* (page 590), and by Kronenberg and Rudensky (page 598). Autoimmune diseases develop when self-reactive lymphocytes escape from tolerance and are activated. Although the mechanisms by which this occurs are not entirely known, autoimmunity is thought to result from a combination of genetic variants, acquired environmental triggers such as infections, and stochastic events.

Table 1 | Simple genetic traits associated with autoimmunity

Gene	Human disease	Mouse mutant or knockout	Mechanism of autoimmunity	References
AIRE	APS-1	Knockout	Decreased expression of self antigens in the thymus, resulting in defective negative selection of self-reactive T cells	2, 3
CTLA4	Association with Graves' disease, type 1 diabetes and others	Knockout	Failure of T cell anergy and reduced activation threshold of self-reactive T cells	9, 65, 66
FOXP3	IPEX (scurfy)	Knockout and mutation	Decreased generation of CD4 ⁺ CD25 ⁺ regulatory T cells	11–13, 67
FAS, FASL	ALPS	<i>lpr/lpr</i> ; <i>gld/gld</i> mutants	Failure of apoptotic death of self-reactive B and T cells	16, 68
C4 complement protein	Associated with SLE	Knockout	Defective clearance of immune complexes and possible failure of B cell tolerance	Reviewed in ref. 69

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Figure 1 | Architecture of single-gene disorders versus a model of autoimmune diseases caused by complex traits. **a**, In simple mendelian traits, the relationship between the causal genetic variant (genotype) and the disease state is deterministic. **b**, In complex traits, the clinically recognized disease state results from interactions between multiple genotypes and the environment. Individual genotypes can affect one or more components of the adaptive or innate immune systems; together these lead to an altered immune response to self antigens. On the basis of current findings, the influence of any individual causal allele is modest, and therefore the relationship between the causal variant and the disease state is probabilistic. Although still providing an incomplete picture, the genetic discoveries in mendelian and common diseases are beginning to help build a model of autoimmune disease. The ultimate goal is to build a specific model for each individual disease whereby the effect of individual risk factors (genetic and non-genetic), their interactions, and their impact on disease susceptibility, disease progression and clinical management, are understood.



Genetics of single-gene disorders

To assess the contribution of genetic factors to disease susceptibility, genetic epidemiologists examine the extent of familial clustering; the degree to which monozygotic twins are more concordant for the presence of a disease compared with dizygotic twins; and the increased risk that family members of persons with disease will develop that disease compared with an individual from the general population. Using such estimates of genetic risk, it becomes obvious that in single-gene disorders, the risk conferred on an individual by a given genetic variant is very high, but the overall impact on the population is minimal because these variants are rare.

In these 'simple' diseases (or traits), although there is often variation in phenotypic expression, the relationship between the causal genetic variant and the disease state is deterministic (Fig. 1a). Genome-wide linkage studies attempt to identify genetic markers (and thus a genomic region) where there is more sharing of alleles between individuals (with a given trait) within families than is statistically expected. Such studies are most commonly used in identifying causal genetic variants for single-gene disorders. Genetic studies of simple traits undoubtedly met with earlier success than those of complex traits. However, recent advances in the study of complex traits are now contributing to this knowledge.

Since inborn errors of metabolism were identified as diseases caused

by mutations or deletions affecting single genes, clinicians and scientists have realized that simple genetic traits can teach us a great deal about the pathways of disease, and about normal physiology. In some cases, the genes identified in simple disorders will have more subtle alterations that confer susceptibility to a 'common' disease (that is, diseases caused by a combination of alleles); in other cases, the mutations will simply suggest which biological pathways may be implicated in common diseases. Although many examples exist of single-gene knockout experimental models that have led to autoimmunity, here we focus on genes that are known to be involved in human disorders (Table 1).

AIRE and central tolerance

AIRE (autoimmune regulator) was identified as the gene that is mutated in autoimmune polyendocrine syndrome (APS-1) — a disorder that manifests as autoimmune attack against multiple endocrine organs, the skin and other tissues². In a *tour de force* of functional genomics, the mouse homologue of the gene has been knocked out, and the *AIRE* protein shown to be responsible for the thymic expression of some antigens that are expressed at high levels in different peripheral tissues. In the absence of thymic expression, T cells specific for these antigens escape negative selection (central tolerance), enter the periphery and attack the target tissues^{3,4}. The possibility that thymic defects underlie autoimmunity was raised

long before this elegant demonstration of the importance of AIRE-dependent thymic selection in self tolerance. For example, one of the susceptibility genes associated with type 1 diabetes in humans is insulin: it has been suggested that the disease-associated polymorphism of the insulin gene reduces its expression in the thymus, thus allowing insulin-reactive T cells to escape deletion⁵. It remains to be determined whether defects in central tolerance contribute significantly to most multigenic autoimmune diseases⁶.

CTLA4 and T-cell anergy

Cytotoxic T lymphocyte antigen 4 (CTLA4; CD152) is an inhibitory receptor expressed by T cells that recognizes the costimulatory molecules B7-1 (CD80) and B7-2 (CD86), the ligation of which shuts off T-cell responses and promotes long-lived anergy⁷. CTLA4 works by competitively blocking the engagement of the activating receptor CD28 (by CD80 or CD86), and by transducing inhibitory signals; the latter probably involves tyrosine and serine/threonine phosphatase activation⁸. Germline *CTLA4* knockout mice develop a fatal syndrome of multi-organ lymphocytic infiltrates and severe enlargement of lymphoid organs. Such symptoms are consistent with a systemic autoimmune reaction presumably directed against multiple, as yet unknown, self antigens. This marked demonstration of the obligatory function of CTLA4 has led to a search for polymorphisms in the *CTLA4* gene that are associated with autoimmune diseases. A surprising discovery was that several such diseases, including Graves' disease, type 1 diabetes and other endocrinopathies, show a striking association with a *CTLA4* polymorphism that results in reduced production of a truncated splice variant, which has inhibitory activity⁹. The functional consequences of producing this altered form of CTLA4 have not been defined, but as expected for a complex disease, the biological effect of the causal allele is more subtle than what is found in monogenic disorders or in knockout mice.

FOXP3 and regulatory T cells

FOXP3 (encoding a transcription factor of the forkhead family) is a striking example of a gene whose role in autoimmunity has been revealed by the confluence of animal studies and studies of a quite rare human disease. CD4⁺CD25⁺ regulatory T cells, now established as major controllers of immune responses to self and other antigens¹⁰, were shown to express high levels of *FOXP3*. Three groups demonstrated that induced knockout or spontaneous mutation of the mouse *Foxp3* gene ('scurfy' mice) led to a systemic autoimmune disease associated with the absence of CD4⁺CD25⁺ regulatory T cells^{11–13}. At the same time, a human disease known by the acronym IPEX (immune dysregulation, polyendocrinopathy, enteropathy, X-linked syndrome) was shown to be associated with mutations in *FOXP3*. These results indicate that the development and/or function of regulatory T cells are dependent on the activity of this one transcription factor; its downstream targets and major functions are not yet defined.

FAS and lymphocyte apoptosis

Fas (CD95) is the prototype of a death receptor of the TNF receptor family. Its biological importance was established by seminal studies

demonstrating that in two mouse models of autoimmunity and lymphoproliferation (named the *lpr/lpr* and *gld/gld* strains), genetic lesions affected the genes encoding Fas and Fas ligand, respectively¹⁴. These results were among the first to show that a single genetic abnormality could give rise to a complex autoimmune phenotype, and that the failure of one apoptosis pathway resulted in self-tolerance breakdown. The Fas death receptor contributes to the deletion of mature T and B cells that recognize self antigens¹⁵. A human disease, called autoimmune lymphoproliferative syndrome (ALPS), which resembles the disease of *lpr* and *gld* mice, is caused by *Fas* mutations¹⁶. Thus, this is an excellent example of how mouse models can lead to a better understanding of human diseases. Although these autoimmune diseases have a superficial similarity to human systemic lupus erythematosus (SLE), there is no convincing evidence for *FAS* polymorphisms or mutations being associated with SLE.

The elegant simplicity of these monogenic disorders has been key to using them to elucidate mechanisms of self tolerance and autoimmunity. The experimental models also illustrate how animal studies inform analyses of human disease, and the striking influences of 'background' genes on the severity and manifestations of diseases. However, as stated at the outset, these single-gene models of autoimmunity are rare; most autoimmune diseases are complex, multigenic traits. The remainder of our discussion will focus on these complex disorders.

Genetics of common autoimmune diseases

In contrast to simple diseases, common diseases are believed to result from a combination of susceptibility alleles at multiple loci, environmental factors (such as smoking, pathogen exposure and hormone levels), and stochastic events (Fig. 1b). There is some debate as to whether common diseases are caused by common alleles of low penetrance (that is, alleles that confer modest increased risk to disease) or by multiple rare alleles of high penetrance. Although most disease alleles identified for common autoimmune diseases have the former characteristics (Table 2), these may not represent the full spectrum of disease alleles. Our ability to discover as yet unidentified disease alleles has been improved by the recent explosion of knowledge regarding the common genetic variation in the human genome. Specifically, recent work has demonstrated that the vast majority of genetic variation in the human genome consists of individual bases that exist as either of two alleles in the population — known as single nucleotide polymorphisms (SNPs) — rather than as deletions or rearrangements. Approximately ten million SNPs in the human genome have a minor allele frequency greater than 1% and represent about 90% of the genetic variation in the human genome. Initial efforts to discover and map SNPs to the reference sequence of the human genome have resulted in a public resource containing most of these common SNPs^{17–19}.

Over the past decade, in autoimmunity as in many other human diseases, there has been great interest in testing candidate genomic regions (identified by linkage studies), or candidate genes (selected on the basis of their location under a linkage peak, or their known functional properties, or both) for evidence of their association with disease. Before the creation of public SNP databases, sequencing was

Table 2 | Relative risk, frequency and population attributable risk for a set of causal variants

Variant	CARD15*	IBD5	PTPN22†	PTPN22†	CTLA4‡	CTLA4‡
Disease	Crohn's disease	Crohn's disease	Type 1 diabetes	Rheumatoid arthritis	Type 1 diabetes	Graves' disease
Relative risk (95% CI)	3	1.3 (1.2–1.4)	1.7 (1.5–1.9)	1.65 (1.2–2.2)	1.14 (1.1–1.2)	1.5 (1.3–1.75)
Frequency	~7%	~35%	~10%	~10%	~50%	~50%
PAR	23%	18%	16%	~12%	13%	36%
References	30	56	37	39	9	9

The genetic characteristics of a selected set of susceptibility genes identified for common autoimmune diseases. The finding of all these loci has been replicated in multiple studies. Therefore, accurate estimates are provided of the risk of disease that each locus confers. Most of these susceptibility loci confer modest risk (1.1–1.7-fold in increased risk to carriers), but given their relatively common frequency in the population (7–50%), have a significant impact on disease burden — as estimated, for example, by the population-attributable risk (PAR). (In contrast, the PAR for causal genes in simple disorders is expected to approach 100%). The modest risk conferred by a given locus in common autoimmune diseases does not provide any significant statistical power to predict whether a given person will develop a disease but may be useful in helping to classify patients into molecularly relevant subgroups. CI, confidence interval; † Calculated for heterozygotes for any of the three characterized mutations (Arg702Tyr, Gly908Arg and Leu1007fsinsC); ‡ Calculated for the 620W allele; § Calculated for the C160/G allele.

needed to identify the genetic variants to test in an association study. Now it seems that there is almost an excess of SNPs to type: do we need to type all ten million to find the few that are causing a given autoimmune disease? To answer this question, it is important to understand the relationships that exist between these SNPs. First, by examining a high density of specific areas of the genome^{20–22}, and by performing genome-wide surveys²³, it has become clear that the alleles of the SNPs form patterns (also known as haplotypes) in the genome. Specifically, the emerging data demonstrate that alleles at nearby SNPs are highly correlated with one another (this is known as linkage disequilibrium). The tight correlation between SNPs, however, is broken down by recombination events resulting in haplotype blocks. Furthermore, the existing data and models based on these data suggest that, rather than occurring randomly, there are recombination hotspots in the genome^{22,24,25}. Knowledge of the haplotype structure not only allows an optimal subset of SNPs to be selected that efficiently extracts the information about the common patterns of variation, but it also can direct how such data should be analysed. For these reasons, an international effort is underway to map the common patterns of genetic variation across the entire genome, and it is expected that the first phase will be completed by mid-2005 (ref. 26). Already the information from this International HapMap can be incorporated into the design of association studies of candidate genes/genomic regions, and it will soon be possible to apply this new information to the design and execution of powerful genome-wide association studies^{27,28}.

Successes in mapping susceptibility genes

Given the early lack of success in identifying genes for complex traits, it was believed that successes in human disease genetics might be limited to disorders with single-gene mendelian inheritance. However, this changed with the publication of several studies reporting the mapping of susceptibility loci in complex human traits: many of these studies took advantage of the haplotype structure of the genome to narrow down the region of association. Early in 2000, fine-scale mapping of a complex human trait successfully led to the localization of the type 1 diabetes locus in the major histocompatibility complex (MHC) to a discrete 570-kilobase (kb) region²⁹. An association mapping approach also proved successful when Hugot and colleagues identified sequence variants in the *NOD2* (*CARD15*) gene that are associated with Crohn's disease³⁰. Comprehensive association mapping approaches successfully identified haplotypic variation in the cytokine gene cluster on chromosome 5q31, which confers susceptibility to Crohn's disease³¹, for which two potentially causal variants have recently been proposed³². The *ADAM33* and *GPRA* susceptibility genes for asthma³³ have similarly been identified. Large association studies of candidate genes have also resulted in important discoveries: the *IDDM1/CTLA4* locus in Graves' disease and in type 1 diabetes⁹; the *NOD2* gene in Crohn's disease^{34,35}; the *PTPN22* gene in type 1 diabetes, rheumatoid arthritis and SLE^{36–39}; and the *PDCD1* gene in SLE and rheumatoid arthritis^{40,41}. The observation that genes such as *CTLA4* and *PTPN22* are associated with multiple disorders is consistent with the hypothesis that certain immunological pathways are common to multiple autoimmune diseases, whereas other pathophysiological mechanisms are specific to a particular disease.

The identification of these susceptibility genes provides the immunology community with an opportunity to study the key pathways and molecular mechanisms that can lead to increased disease susceptibility. Initial functional studies have begun to provide some relevant clues. For example, work regarding the effect of genetic variants in *CTLA4* on susceptibility to type 1 diabetes in mice and humans has provided substantial evidence that modulating the expression of alternative splice forms can regulate the expression of T-cell-mediated autoimmunity^{34,42}. Specifically, the human variant leads to the decreased production of a ligand-independent form of *CTLA4* — a splice form believed to be involved in the downregulation of memory/effector T-cell activation^{34,42}. Genetic variation in the human *PTPN22* gene also seems to modulate T-cell activity. However, this is

due to an amino acid change in the encoded protein, the lymphoid protein tyrosine phosphatase (LYP), a known suppressor of T-cell activation. Bottini and colleagues⁴³ hypothesized that *PTPN22* was a good candidate gene for autoimmunity because it and other protein tyrosine phosphatases are involved in preventing spontaneous T-cell activation. These investigators identified an SNP that was associated with type 1 diabetes⁴³, which changed an amino acid involved in the interaction between LYP and the negative regulatory kinase CSK. *PDCD1*, a positional and functional candidate gene, encodes the programmed cell death 1 gene whose gene product has been shown to regulate peripheral tolerance in T and B cells^{44,45}. An intronic variant in the *PDCD1* gene was proposed to lead to aberrant *PDCD1* regulation and increase an individual's susceptibility to SLE⁴¹. These few examples demonstrate how variants that modify gene expression or protein structure can lead to modulation of the adaptive immune responses.

Alteration of the innate immune system, on the other hand, may be the primary effect of the two missense mutations and the truncation mutant in the *NOD2* gene, all of which have been associated with Crohn's disease. Specifically, the *NOD2* gene product has been shown to recognize muramyl dipeptide (MDP) — a component of the bacterial wall. The Crohn's disease-associated mutations were believed to lead to impaired nuclear factor- κ B (NF- κ B) signalling following recognition of MDP^{34,46,47}. Using spleen macrophages derived from *Nod2*-deficient mice, Watanabe and colleagues proposed an alternative explanation. Specifically, they suggested that the *NOD2* gene product limits the pro-inflammatory effects mediated by Toll-like receptor 2 (TLR2) signalling, such that mutations in *NOD2* would lead to excessive T-helper type 1 (T_H1) responses^{46,47}. Efforts to knock out the *Nod2* gene have not, however, led to gastrointestinal inflammation. This perhaps is not surprising given the complexity of the human disease; no single genetic or environmental factor is expected to be necessary or sufficient to cause the disease. These *Nod2*^{-/-} mice also did not show differential susceptibility to chemically induced gastrointestinal inflammation, but they did seem to be more susceptible to oral challenge with Gram-positive bacteria^{48,49}. The results from knocking-in the human frameshift mutation into mice (*Nod2*^{2939C}), however, seem to suggest a very different mechanism: as opposed to the lack of response to MDP in the *Nod2*^{-/-} mice, the *Nod2*^{2939C} mice have an elevated response to MDP and a heightened intestinal inflammatory response to chemical injury⁵⁰.

These exciting experiments demonstrate the importance of a genetic and environmental context for such functional studies and the challenges that lie ahead in deciphering the exact mechanisms by which genetic variation can lead to autoimmunity. Although it is still too early to know the precise mechanisms by which individual allelic variants confer susceptibility, the elucidation of the pathogenic mechanisms in the coming years will undoubtedly shed much light on the common and disease-specific processes.

Challenges in mapping susceptibility genes

Genetic association studies of autoimmune diseases, as for all other complex human traits, face two main challenges: the ability to distinguish between true and false associations, and the ability to demonstrate causality. The first challenge relates to the nature of many causal alleles in common disease; many of the implicated alleles only confer a modest increased risk of developing disease (see Table 2). The power to detect disease susceptibility genes is influenced by the magnitude of the risk conferred by the susceptibility allele, and by its frequency in the population. The weaker the effect of an allele, the greater the sample size required to detect an association signal that can be distinguished from the background noise of an association study. Unfortunately, it is not possible to know *a priori* the frequency and the strength of disease alleles. However, the detection of modest alleles may require many thousands of samples²⁸. This is further complicated by the fact that the first report of an association often overestimates the effect of the putative associated allele⁵¹. This effect is known as the 'winner's curse' — a mathematical concept first described in the context of auctions and bilateral negotiations⁵². Replication studies, there-

fore, often produce seemingly inconsistent results when compared with the original study^{53,54}. Luckily, this challenge can be addressed by performing association studies (novel and replication) with large sample sizes, and by combining the results from multiple studies using a meta-analysis approach^{53,55}. Indeed, these two approaches have successfully demonstrated that a number of susceptibility genes for autoimmune disease (for example, *PTPN22* in type 1 diabetes, *IBDS* and *NOD2* in Crohn's disease) can be convincingly identified and replicated^{37,56,57}.

The second challenge relates to the patterns of linkage disequilibrium described above. Although intervals of historical recombination occur between haplotype blocks, this is a modest enough effect for correlations between blocks to persist²¹. Given the extreme local variation in the magnitude of this recombination²⁵ and in gene density¹⁹, genomic regions implicated by association studies may contain a single gene or multiple genes. For the latter, current sample sizes may be insufficient to narrow down the implicated region to less than a handful of genes. One example is a 250-kb haplotype on chromosome 5q31 that was found to be associated with Crohn's disease³¹. The haplotype associated with increased risk of developing Crohn's disease extends uninterrupted across multiple blocks, as the recombination events that have occurred in the intervals in this region have primarily involved the haplotypes that do not confer risk. Therefore, this association implicates a region that contains a minimum of four genes (*IRF1*, *SLC22A5* (*OCTN2*), *SLC22A4* (*OCTN1*) and *PDLIM3*) and multiple SNPs that have an allele that is unique to the risk haplotype. Therefore, the identification of the causal gene(s) and related variant(s) will rely on functional studies that link gene/variants and phenotype. Another important example is that of the MHC, where extended haplotypes are believed to represent a common feature of this region⁵⁸. Importantly, the MHC region has been associated with almost all autoimmune diseases and contains the HLA genes: HLA gene products are crucial for antigen presentation to cells of the adaptive immune system and for controlling some reactions of the innate immune system, such as activation of natural killer (NK) cells. This genomic region also contains hundreds of other genes, many with putative or proven function in the immune system⁵⁹. Most association studies of this genomic region, however, have been limited to one or a few of the HLA genes. The preliminary efforts to map the patterns of genetic variation in this region suggest that dense sets of SNPs applied to large study cohorts will enable the identification of the HLA and non-HLA components of the genetic susceptibility conferred by the MHC region⁶⁰⁻⁶³.

Future prospects

One of the great promises of genetics is to help build a molecular model of disease that combines information regarding genotype and environment, as well as their various interactions (Fig. 1b). One challenge that is specific to immune-related disorders is the fact that an individual's functional repertoire of specific antigen receptors (B-cell receptors and T-cell receptors) is a product of the interaction between their genetic repertoire and the environment. It will therefore be important to define an individual's functional state in the context of their genetic background. Despite this challenge, some of the discoveries so far in single-gene and complex forms of autoimmunity are helping to build such a model. These models could potentially help to assess an individual's risk of developing disease. There is also some indication that this information will be relevant for assigning patients to molecular subgroups of these very heterogeneous diseases, which may help to predict a particular disease outcome and/or response to therapy.

A complementary approach to improving the response to current therapies is to examine variation in the genes controlling the absorption, distribution, metabolism and excretion of the drugs used to dampen the chronic inflammatory response in patients⁶⁴. An aspect of these models that may prove to be important is knowing which genes are common to more than one autoimmune disease versus those that are disease-specific, as these might turn out to be qualitatively very different drug targets. Finally, there is no doubt that any novel suscepti-

bility gene, or even newly identified causal variation in a previously well characterized gene, will provide the immunologist with some very interesting avenues to follow in their path to understanding the fundamental mechanisms of autoimmunity.

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Acknowledgements We thank J. Bluestone, M. Anderson, M. Daly and F. Sanchez for their helpful comments. We also thank L. Gaffney for her help in the preparation of the manuscript. J.D.R. is supported by grants from the CCFA and NIDDK.

Competing interests statement The authors declare that they have no competing financial interests.