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Richardson, Naomi; Wraith, David Cameron

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Review

Advancement of antigen-specific immunotherapy: knowledge transfer between allergy and autoimmunity

Naomi Richardson^o and David Cameron Wraith^{*}

Institute of Immunology and Immunotherapy, College of Medical and Dental Sciences, University of Birmingham, Birmingham B15 2TT, UK

^{*}Correspondence: David Cameron Wraith, Institute of Immunology and Immunotherapy, College of Medical and Dental Sciences, University of Birmingham, Birmingham B15 2TT, UK. Email: D.Wraith@bham.ac.uk

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Summary

Targeted restoration of immunological tolerance to self-antigens or innocuous environmental allergens represents the ultimate aim of treatment options in autoimmune and allergic disease. Antigen-specific immunotherapy (ASI) is the only intervention that has proven disease-modifying efficacy as evidenced by induction of long-term remission in a number of allergic conditions. Mounting evidence is now indicating that specific targeting of pathogenic T cells in autoinflammatory and autoimmune settings enables effective restoration of immune homeostasis between effector and regulatory cells and alters the immunological course of disease. Here, we discuss the key lessons learned during the development of antigen-specific immunotherapies and how these can be applied to inform future interventions. Armed with this knowledge and current high-throughput technology to track immune cell phenotype and function, it may no longer be a matter of ‘if’ but ‘when’ this ultimate aim of targeted tolerance restoration is realised.

Keywords: immunotherapy, immune tolerance, allergy, autoimmunity, immunoregulation

Introduction

The treatment of allergy and autoimmunity urgently requires novel therapeutic approaches; current medical interventions broadly aim to manage symptoms of

disease but do not address their underlying cause, i.e. loss of immunological tolerance. Immunosuppressive drugs have both short- and long-term adverse effects, most importantly compromised immune function in immune

Abbreviations: AIT: Allergen immunotherapy; APC: Antigen-presenting cells; ASI: Antigen-specific immunotherapy; BCR: B cell receptors; Breg: Regulatory B cells; EAE: Experimental autoimmune encephalomyelitis; LSEC: Liver sinusoidal endothelial cells; MBP: Myelin basic protein; MHC-II-NP: MHC class II conjugated nanoparticles; moDC: Monocyte-derived DC; MS: Multiple sclerosis; NP: Nanoparticles; PIT: Peptide immunotherapy; SCIT: Subcutaneous immunotherapy; SLIT: Sublingual immunotherapy; ssDC: Steady-state DC; TCR: T cell receptor; Teff: Effector T cell; Tr1-like: Type 1 regulatory-like; Treg: Regulatory T cells; TSHR: Thyroid-stimulating hormone receptor.

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antigen. Genetically modified recombinant antigens have been designed with mutated IgE-binding motifs or as fragmented constitutive overlapping peptides to reduce the risk of IgE cross-linking, while maintaining T cell reactivity and represent a powerful tool for engineering a safer product for desensitisation [27–29].

The complexity of autoimmune diseases poses a significant challenge to antigen identification. Immune responses vary considerably between patients and at different time points of disease progression [30, 31]. At present, our knowledge of disease-initiating and propagating autoantigens in many autoimmune diseases is incomplete and further complicated by epitope spreading [32]. Despite this, ASI has shown promise in inducing tolerance towards specific auto-antigens. A series of studies in the 1980–90's indicated that disease in rodent models of autoimmune disease including experimental autoimmune encephalomyelitis (EAE) [33, 34], collagen-induced arthritis [35], and non-obese-diabetes [36] could be ameliorated by ASI. More recently, clinical trials utilising tolerogenic peptides in the treatment of multiple sclerosis (MS), type 1 diabetes, systemic lupus erythematosus, and Graves' disease have been safe, well tolerated and indicate that disease severity can be lessened [37–39]. Such trials are the outcome of decades of research into the identification of relevant auto-antigens and T cell epitopes in these diseases.

Experience has shown that when the pathogenic autoantigen is defined, e.g. thyroid-stimulating hormone receptor (TSHR) in Graves' disease, it is possible to target disease pathogenesis and deliver clinical benefit [40]. Where the autoantigen(s) responsible are not fully defined or disease is driven by reactivity to multiple antigens, it is possible to control disease severity by targeting only one antigen within the same affected tissue via bystander or linked suppression.

Immune regulation: the need for active suppression and bystander regulation

Linked suppression occurs when antigen-specific T cell tolerance induction to an immunodominant epitope of antigen A leads to suppression of immune responses against other epitopes within antigen A. Bystander suppression enables antigen-specific T cells directed against antigen A to indirectly dampen immune responses against antigens B, C, and so on, by involvement of T cell-mediated suppression of antigen presenting cells and neighbouring T cells (Fig. 1). Both linked and bystander suppression have been reported outcomes of ASI in multiple allergic and autoimmune disease settings.

The processes by which this localised antigen-independent suppression occurs are still poorly understood, although bystander suppression plays an identifiable role in murine peptide tolerance models of EAE and in allergic contexts [41, 42]. In cat allergy, tolerance induction using 12 Fel d1 peptides not only suppressed patient responses to these Fel d1 peptides, but also to Fel d1 peptides not included in the therapy [43].

IL-10, secreted by anergic Type 1 regulatory-like (Tr1-like) cells, regulatory T cells (Treg), regulatory B cells (Breg), and tolerogenic dendritic cells, is thought to be central in establishing broader regulation following antigen-specific therapy [44]. Its role in establishing bystander suppression is likely due to its ability to downregulate costimulatory molecules and MHC-II on the surface of antigen-presenting cells (APC) [45–47], thus reducing antigen-presentation and T cell priming potency of APC. IL-10 is also able to directly suppress both T and B cell responses via inhibition of co-stimulatory signalling [48–50]. This not only suppresses subsequent immune responses to the initial antigen targeted, but also other disease-relevant antigens nearby in the inflamed tissue.

Tolerance-induced Tr1-like and Treg express high levels of coinhibitory receptors CTLA-4, LAG-3, PD-1, TIM-3, and TIGIT [51, 52]. The inhibitory receptors control T cell signalling through mechanisms including competition with ligands/counter receptors, engagement of protein phosphatases and inhibitory signalling. Collectively, they act as checkpoints and fine tune the magnitude of the T cell response to antigen [53].

TGF- β is highly expressed by Treg as a result of oral antigen delivery [54] and contributes to prevention of EAE when disease is initiated via myelin basic protein (MBP) or proteolipid protein – indicating strong bystander control of multiple antigen specificities in complex disease [55]. Targeting antigens to the liver induces Treg in a TGF- β -dependent manner [56] and has also been shown to generate multi-antigen tolerance induction [57].

Antigen-specific immunotherapies based on single antigen specificities are unlikely to be effective in complex and dynamic multi-antigen diseases such as type 1 diabetes and rheumatoid arthritis, unless they can evoke bystander suppression [58]. Therefore, understanding the mechanism of bystander suppression and how best to incorporate it into antigen-specific immunotherapy will prove crucial to resolve the dilemma of which antigen(s) to target in a specific disease. Providing that tolerance induction towards a dominant antigen is sufficient to control the pathogenic nature of T cells of multiple antigen specificities, disease severity should be ameliorated.

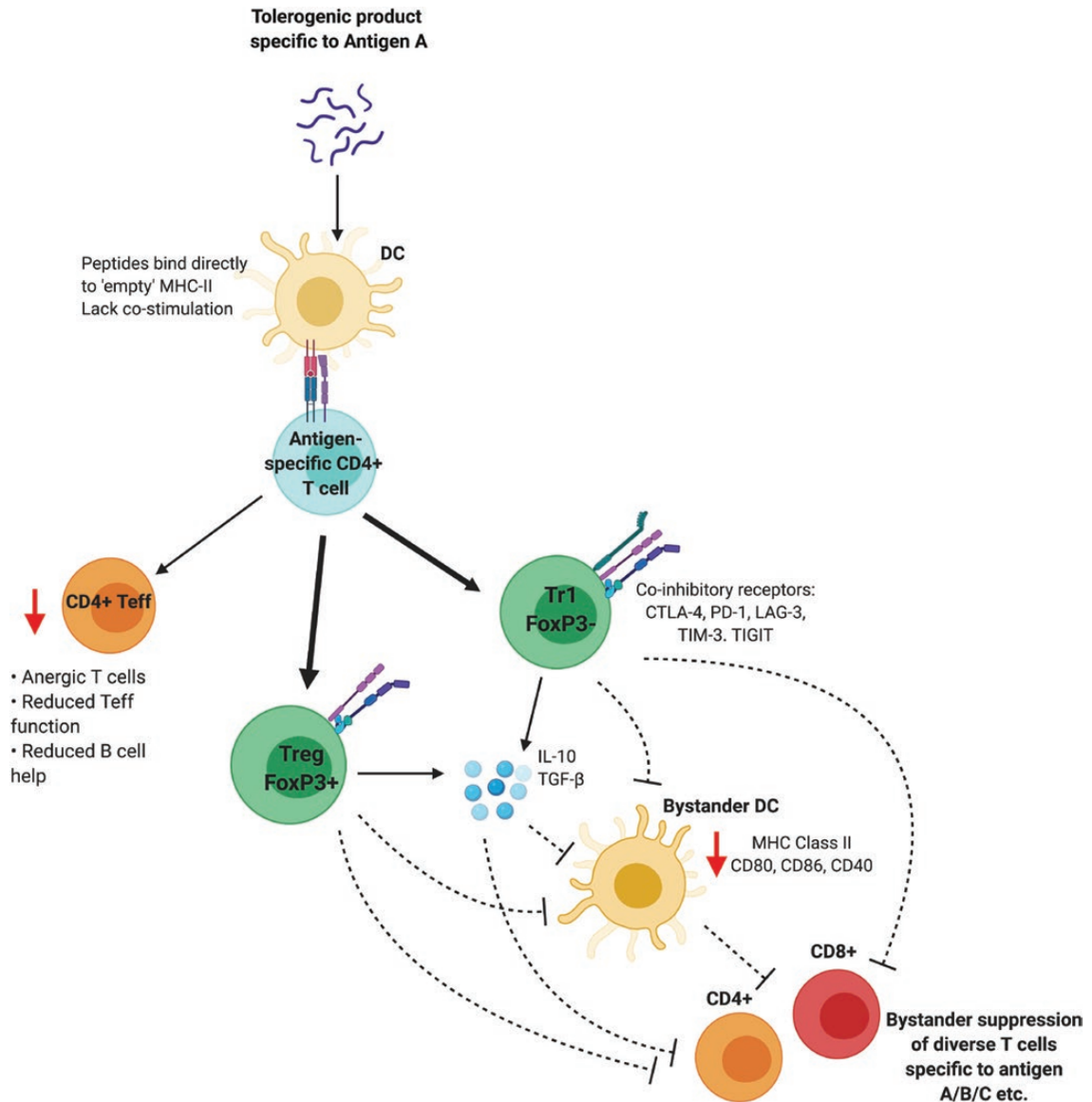


Figure 1 Proposed mechanism of action of bystander suppression. Antigen-specific immunotherapies prevent the generation and activation of CD4⁺ Teff and instead divert Tconv CD4⁺ cells towards anergy and also promote the expansion of antigen-specific Tr1-like cells and/or Treg. Both tolerised Tr1-like and Treg can exert cell-contact mediated and cytokine mediated suppression (dashed lines) on APC and non-antigen-specific T cells to ultimately prevent T cell activation in a non-antigen-specific manner.

Mechanism of action and associated risks

Through careful investigation of ASI/AIT using either intact allergen, autoantigen, or antigenic peptides, we now have a good understanding of the cellular and molecular mechanisms involved in tolerogenic antigen delivery and the risks associated with each type of approach (summarised in Fig. 2).

Allergen immunotherapy using intact allergen commonly results in a decrease of allergen-specific effector T cell (Teff) number and/or functionality, often described as a Th2→Th1 population shift, although we would argue this is often related to a change in ratio between these populations as opposed to Th2 converting to Th1 [8, 59–61]. Regulatory populations are elevated after

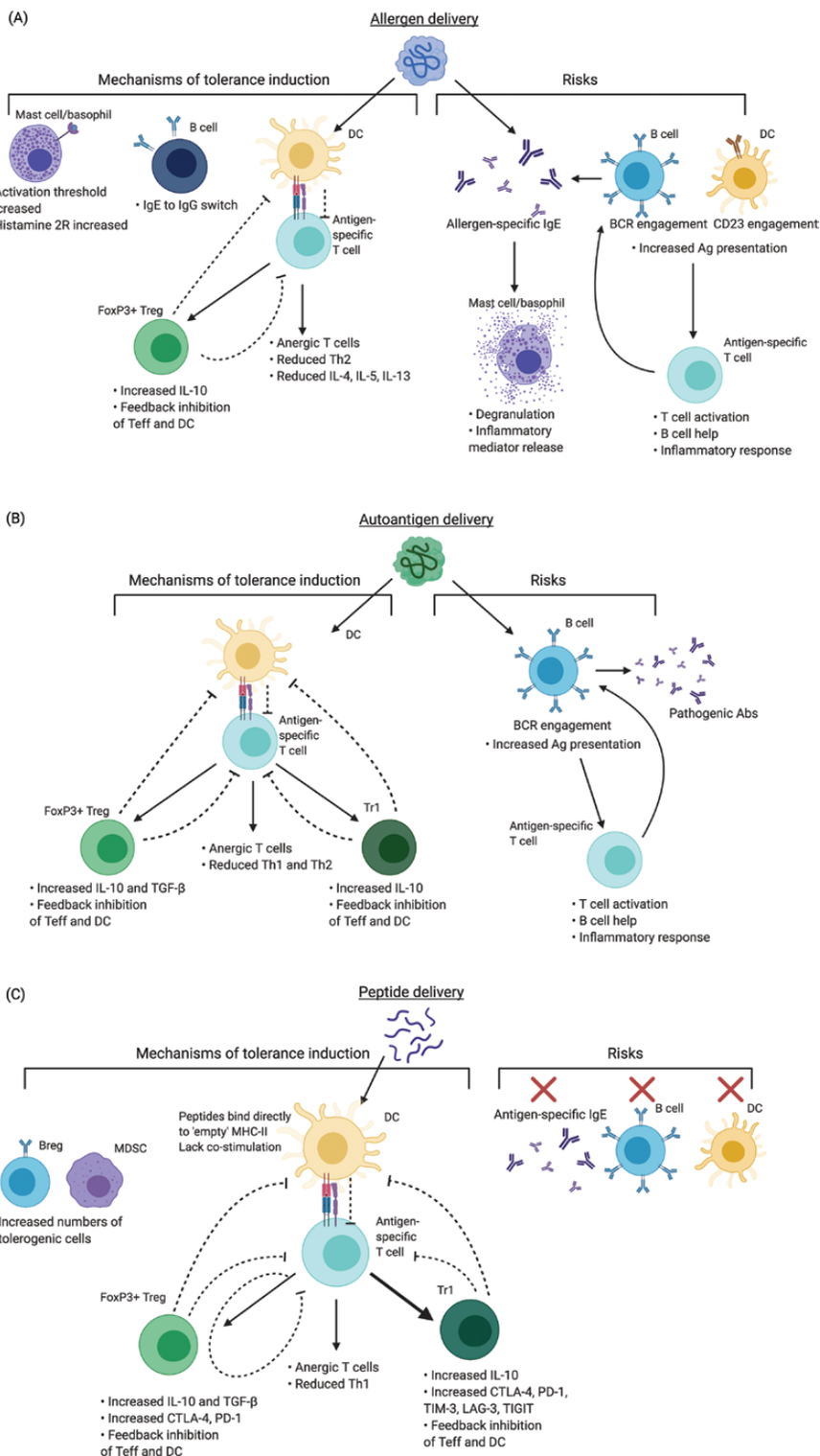


Figure 2 Summarised mechanisms of action of ASI/AIT and associated risks. Antigen-specific immunotherapies have varying mechanisms of action and potential risks depending on whether they utilise (A) intact allergen, (B) intact autoantigen, or (C) peptides representing T cell epitopes of either allergen or autoantigen. Promotion of activity denoted by black arrows, inhibition of activity denoted by black dashed lines and mitigation of risks denoted by red crosses.

treatment; some studies report a dominant FoxP3⁺ Treg effect while others report primarily FoxP3-like [8, 62]. This may be disease-specific, related to the nature of antigen delivered, treatment protocol used, or simply the design of immunological readouts. The consensus, however, is that peripherally induced regulatory T cells are expanded after treatment and contribute to disease control. AIT also moderates basophil and mast cell degranulation, increasing the threshold required for their activation, in addition to increasing expression of histamine receptor 2 to act as a histamine 'sink' [17, 63]. IL-10⁺ are promoted by AIT intervention [64–66]. Most importantly, allergen-sequestering IgG4 titres are increased. IgG4 competes directly with IgE for antigen-binding sites, reducing the likelihood of early phase immune response on subsequent exposure [13, 66–69]. Of the 4 IgG subclasses, IgG4 has the lowest abundance, accounting for around 4% of serum IgG, yet it can reach up to 75% IgG after AIT [69]. IgG4 has several 'anti-inflammatory' properties, due to its low affinity for Fcγ receptors, inability to activate complement and ability to form bivalent antibodies which are not able to cross-link antigen to form immune complexes [70, 71]. The production of IgG4 is mediated by plasmablasts/plasma cells [72, 73] differentiated from IL-10⁺, where IL-10 promotes the generation of 'blocking' IgG4 antibodies, while inhibiting IL-4-mediated IgE class-switching in humans [17, 74, 75]. Although the IgG4-mediated suppression of IgE is well documented, more recently, evidence has emerged suggesting that antibodies of different classes, particularly IgG2, can also play a role in blocking IgE engagement [73, 76, 77]. Decrease in allergen-specific IgE has been observed after long-term treatment duration (1–2 years), occurring much later than symptomatic relief [17].

Conversely, there are well-documented risks associated with use of whole allergen: even very low doses of pure antigen can cause unpredictable cross-linking of IgE and activation of mast cells and basophils via the high-affinity IgE receptor FcεR1. IgE-antigen complex bound by the lower affinity IgE receptor, CD23, on B cells and DC promotes antigen uptake and efficient presentation to T cells [78, 79], perpetuating allergen-specific IgE production, T cell priming, and activation. Furthermore, conformational epitopes of antigen can directly bind B cell receptors (BCR) for BCR cross-linking [80].

ASI directed towards autoimmune diseases also initially used whole autoantigen as the tolerising agent. Early trials in MS injected intact MBP isolated from human, porcine, or bovine sources and did not promote immunological or symptomatic improvement [81–83]. The delivery of whole antigen proved to be high risk,

due to the potential generation of pathogenic antibodies [84]. As such, considerable progress was made to properly identify relevant T cell epitopes in murine models and MS patients [58, 85–87] for use in peptide immunotherapy (PIT).

Peptides representing T cell epitopes have also been employed in the allergy field, as peptides avoid IgE-mediated immune responses and unpredictable immunological effects associated with the use of whole allergen [88]. Short soluble peptide epitopes are unable to cross link IgE and are unlikely to provide the 3D-conformation required to function as B cell epitopes. Peptides are significantly less likely to result in mast cell and basophil degranulation compared to whole allergen [22, 89]. The mechanisms of tolerance induction when utilising whole allergen versus peptide-based approaches, are likely to be subtly different, although direct mechanistic comparison studies between the two parallel approaches are lacking at present. Akdis and colleagues showed that peptide immunotherapy did not generate B cell tolerance – one of the key features reported via use of whole allergen in AIT. However, these experiments did generate 'blocking' IgG4 antibodies and a relative reduction in IgE [13, 14].

Where whole antigen requires processing by APC for presentation to T cells, peptides representing disease-relevant T cell epitopes specifically utilise resting DC in lymphoid organs for presentation to cognate T cells without the need for antigen processing [90]. Steady-state DC (ssDC) are tolerogenic and well-suited to promote the restoration of T_H17 versus Treg balance. A proportion of MHC Class II on ssDCs are 'empty' or transiently loaded with low-affinity peptides [91]; therefore, exogenous peptides delivered can bind directly to MHC-II for presentation to CD4⁺ T cells. ssDC provide low levels of costimulation (CD80/CD86) to T cells and are less efficient in antigen uptake and presentation [92, 93]. As such, antigen-specific T cells do not receive sufficient stimulatory signal from T cell receptor (TCR) engagement alone to become activated [94] and are instead diverted into a state of functional anergy [95] by repeated antigen exposure in which they no longer respond to antigen via classical inflammatory signalling pathways but instead exert a regulatory phenotype. Antigen-specific naive and effector CD4⁺ T cells become regulatory Tr1-like cells (FoxP3⁻) and FoxP3⁺ Treg throughout PIT [96] and express high levels of IL-10 and co-inhibitory receptors (CTLA-4, PD-1, TIGIT, LAG-3) [52]. As a result, T cell immunity directed towards the antigen is quenched; readouts often include significant reduction in T_H17 cytokine production (IFN-γ, IL-2 in autoimmunity; IL-4, IL-5 in allergy) [97].

Peptide design must reflect naturally processed T cell epitopes, with high solubility and minimal aggregate potential. Studies in MS using an altered peptide ligand warned the field that using non-native peptides could result in disease exacerbation [98, 99]. These adverse effects primarily arose due to administration of an excessively high dose of peptide which may not have remained soluble *in vivo*, hence promoting rather than suppressing immunity. This story highlights the need for peptides used in antigen-targeting immunotherapies to be highly soluble and to mimic the naturally processed T cell epitope to avoid unforeseen immunological consequences. These risks were avoided in later clinical trials utilising natural T cell epitope peptides with high solubility [37, 100].

Route of administration

Tolerance induction via mucosal surfaces (oral, nasal, sublingual) has been popular historically, as these sites are exposed continually to environmental antigens and yet in healthy individuals do not generate immune responses to these stimuli.

Seminal experiments pioneered by Weiner and colleagues in a number of animal autoimmune disease models, showed overwhelming efficacy of fed antigen to prevent disease [53]. Oral tolerance was notably less effective in pre-sensitised animals (which better reflect ongoing disease in humans) [101]. Unfortunately, in clinical trials, oral tolerance induction in MS using MBP was deemed to be safe but ineffective. This is most likely due to the relative low doses of antigen used in patients compared to those tested in animals [102] and to generally 'weak' immune responses towards autoantigens.

Even in allergic diseases where the antigen typically generates stronger immune responses, oral delivery of antigen does not consistently achieve tolerance. An exception to this is peanut allergy, in which repeated doses of pure peanut protein increasing up to 800 mg were shown to decrease peanut sensitivity after 30 weeks of treatment. Patients were not followed up after treatment had ended, therefore the longevity of reduced sensitivity and the requirement for maintenance therapy was not assessed [103]. Delivery of the offending antigen to the site of hypersensitivity may co-opt natural regulatory feedback loops *in situ* for disease modification. Such a significant amount of protein would be extremely expensive when requiring recombinant allergens, and highly inefficient due to degradation within the stomach prior to having any tolerogenic effect in the gut.

Mucosal delivery via sublingual immunotherapy (SLIT) and systemic delivery via subcutaneous immunotherapy (SCIT) routes offer clinical efficacy using much

lower doses of antigen and are now common practice in allergen immunotherapy [11, 12]. Few studies compare the efficacy of SCIT versus SLIT directly, making an over-arching judgement on the validity of each method difficult; however, the mechanism of action is likely to be subtly different [104, 105].

Intralymphatic antigen delivery is early in development, but has shown remarkable efficacy in murine models [106] and in clinical trials of allergy [107, 108]. Direct delivery of grass pollen allergen intralymphatically has generated safe, pain-free, and effective allergen-specific tolerance much more rapidly than standard SCIT therapy (8 weeks with 3 injections vs. 3 years therapy with 54 injections). Allergy symptoms and allergen-specific IgE were significantly reduced after both treatment courses and maintained for 2 years post-treatment. It is likely that this approach is transferable across allergies, upcoming trials will be followed with interest.

In the context of autoimmune disease, thorough pre-clinical investigation in mouse models of disease have shown a hierarchy of delivery route efficacy, with subcutaneous > intranasal > oral delivery [109]. As such, clinical trials in relapsing remitting MS and Graves' disease were performed by subcutaneous/intradermal delivery of tolerogenic peptides. No unexpected safety concerns arose during these trials, and both displayed significant decreases in disease severity by the end of treatment course [37, 110]. Importantly, studies in experimental animal models have shown that s.c. injection of soluble peptides are detected on the surface of ssDC within minutes [90]. Naive T cell encounter with the epitope presenting ssDC transiently signal via their TCR, as evidenced by ERK phosphorylation followed by transient IL-2 secretion; however, both ERK phosphorylation and inflammatory cytokine secretion are reduced with further antigen administration. Repeated delivery of soluble peptide leads to induction of IL-10 expression in the anergic T cells [109, 111].

The application of ASI via the intralymphatic route (DIAGNODE trial) in autoimmune disease used direct injection of glutamic acid decarboxylase antigen into lymph nodes of type 1 diabetes patients, with a promising reduction in insulin requirement after treatment [112, 113]. This alteration in delivery route may be a more potent means of generating immune tolerance, as suggested by murine and allergy studies; however, this approach is less practical for tolerance maintenance.

Dosing strategy and longevity of response

Dose escalation has been a cornerstone of allergen immunotherapy ever since Freeman and Noon's very first

NPs are taken up by different APC depending on their size. Small NP are endocytosed by DC; Kishimoto and colleagues have delivered rapamycin to DC with antigen in order to induce regulatory T cells [127]. Larger NP containing antigen is phagocytosed by macrophages in order to create a suppressive immune response [128]. Preclinical work describing encapsulation of gliadin [129] has led to a clinical trial of gliadin NP in coeliac disease. Santamaria and colleagues have described a sophisticated NP delivery approach. Here NP are coupled to MHC class II molecules and incubated with peptide epitopes. These MHC-II-NP do not activate naive T cells but promote IL-10 production by antigen-specific Th1 cells [130]. The induction of Tr1-like cells by MHC-II-NP was recently shown to mediate bystander suppression of autoimmune responses in the liver [57, 131].

How best to deliver antigens for tolerance induction

- a. Is it necessary to couple antigens to NP for tolerance induction? The use of NP arose from early studies in which it was shown that peptide epitopes can induce an allergic response *in vivo* [132]. In our experience, however, the balance between a peptide epitope being tolerogenic rather than immunogenic is determined by its solubility. Furthermore, peptides themselves directly target tolerogenic DC *in vivo* when designed to mimic naturally processed antigens. Our original observations showed that some but not all T cell epitopes induce tolerance when administered in a soluble form [133]. Peptides must be designed to bind MHC II in a conformation that mimics the naturally processed epitope in order to induce tolerance. This is consistent with our recent observation that tolerogenic peptides bind directly to steady state DC *in vivo*. DCs collected from lymphoid tissues following subcutaneous injection of soluble peptide are able to induce tolerance following adoptive transfer in mice [90]. Furthermore, insoluble peptides fail to reach lymphoid DC following subcutaneous injection and are immunogenic rather than tolerogenic. However, these peptides are rendered tolerogenic by increasing their solubility. The first rule governing design of peptides for tolerance induction is, therefore, peptides must mimic naturally processed epitopes when bound to their MHC restriction element.
- b. Peptides must be soluble such that they rapidly distribute throughout the body and bind to MHC II on ssDC in lymphoid organs.
- c. Peptides should induce cytokines that promote bystander suppression such that an epitope from antigen
- d. Peptides with the properties listed above are defined as antigen processing independent epitopes or apitopes.

ASI using tolerogenic peptides: mechanism of action and translation to the clinic

Our recent work has defined the detailed mechanism of how tolerogenic peptides function *in vivo*. Our original work compared mucosal routes of administration. Oral delivery of peptides was ineffective due to proteolytic destruction [116] whereas nasal administration induced bystander suppression in a dose dependent fashion [9, 45, 134]. Peptide therapy induced cells with a Tr1-like, IL-10 secreting phenotype [135] that mediated suppression by downregulating the antigen presenting properties of DCs [136]. The mechanism by which soluble, tolerogenic peptides convert potentially pathogenic T cells into Tr1-like cells was revealed in recent studies. First, Burton *et al.* showed that repeated encounter with peptides presented by ssDC induced antigen-specific CD4 T cell anergy and suppressed secretion of inflammatory cytokines [52]. Analysis of gene expression in cells showed that peptide treatment caused a marked upregulation in expression of genes encoding inhibitory receptors PD1, CTLA4, Lag3, Tim3, and TIGIT and transcription factors known to promote expression of IL-10 such as c-Maf. This transcriptional signature was also seen in other Tr1-like cells and in tumour infiltrating lymphocytes [137]. Later, our work has revealed the link between antigen-exposure, T cell signalling, and the subsequent expression of IL-10 and the generation of Tr1-like cells. The anergy seen among T cells in peptide-induced tolerance results from a membrane proximal block in cell signalling causing a loss of inflammatory cytokine gene expression [95]. Bevington *et al.* have shown that this reduced level of cell signalling is insufficient to drive the epigenetic changes required for transcription of inflammatory genes; however, epigenetic priming of genes associated with tolerance renders them sensitive to reduced levels of transcription factors [111]. This novel mechanism explains how cells including tumour infiltrating lymphocytes and cells rendered tolerant with either peptide antigens or anti-CD3 antibodies [138] change their transcriptional landscape with selective upregulation of genes encoding inhibitory receptors, transcription factors

such as c-Maf and the anti-inflammatory cytokine IL-10. Furthermore, the detailed understanding of how tolerogenic peptides modulate the immune response to antigen provides the foundation for their application in treatment of hypersensitivity diseases including auto-immune and allergic diseases.

Antigen-specific immunotherapy with apitopes has been tested in four clinical trials in two autoimmune diseases with distinct immune pathologies. Multiple sclerosis is a cell-mediated disease with various disease-associated antigens. Two phase 1 followed by a phase 2 clinical trials have shown that treatment with a cocktail of four HLA-DR2 binding peptides from MBP_{Ac1-9} was sufficient to significantly suppress inflammation in the CNS as measured by gadolinium enhanced MRI [37, 100] and to improve cognition in patients with relapsing MS. In Graves' disease autoimmunity is caused by antibodies specific for TSHR. Two dominant HLA-DR3 binding peptides suppressed immune responses in HLA-DR transgenic mice [139]. Furthermore, intradermal injection of these peptides normalised thyroid hormone secretion in 7/10 patients with mild-to-moderate hyperthyroidism in a phase 1 trial. Most importantly, the results of these four clinical trials show that treatment with soluble peptides designed as apitopes is well tolerated with promising signs of efficacy. It is important to add that these clinical trials used a dose-escalation protocol shown to promote Tr1-like cell generation in pre-clinical models. Recent studies with peptide immunotherapy in coeliac disease have proved the importance of dose escalation [139]. The dose-escalation protocol shown to induce Tr1-like cells through epigenetic modification of the genome in experimental animal models [111] has proved to be the preferred approach for effective tolerance induction in the clinic. Further analysis of antigen-specific T cells in future clinical trials of antigen-specific immunotherapy is required to confirm that this is due to selective epigenetic priming at tolerance-associated genes.

Concluding statement

Antigen-specific immunotherapy remains the 'holy-grail' for selective treatment of allergies and autoimmune diseases. Rapid advances in our understanding of the mechanisms involved provide options ranging from the administration of tolerogenic DC, through design of sophisticated NP to simple delivery of apitopes. Critical issues including mechanism of action, bystander suppression, ease of manufacture, and successful translation to the clinic will determine success of each approach for treatment of hypersensitivity diseases.

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Author contributions

Authors listed have made a substantial, direct, and intellectual contribution to the work, and approved it for publication. Authors are accountable for all aspects of accuracy and integrity of the work.

Conflict of interest

D.C.W. is Professor of Immunology at the University of Birmingham and CSO and Founder of Apitope International NV. N.R. declares that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Data availability

The data underlying this article are cited in the reference list and available in the public domain.

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