



Original Article

Modulating Immune Checkpoints: A Promising Approach for Personalized Therapy in Autoimmune Disorders

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ABSTRACT

The modulation of immune checkpoints has emerged as a compelling therapeutic strategy for managing autoimmune disorders, offering a targeted alternative to conventional immunosuppressive therapies. This study explores the evolving role of checkpoint inhibitors such as CTLA-4, PD-1, PD-L1, and novel targets like OX40 and LAG-3 in restoring immune tolerance while minimizing systemic side effects. Through a comprehensive secondary analysis of recent clinical trials and preclinical investigations, the research identifies key therapeutic outcomes associated with checkpoint inhibition, including reductions in disease flare frequency, improved biomarker profiles, and enhanced remission rates in conditions like rheumatoid arthritis, systemic lupus erythematosus, and multiple sclerosis. The results presented in 4 tables demonstrate successfully running trials, the need for biomarker-based patient subtyping, and the effectiveness of unique strategies like CAR-T cell therapy and engineered regulatory T cells to eliminate autoreactive immune cells selectively. Furthermore, eleven figures that are presented additionally emphasize the effects of these therapies on outcomes, providing information about immunomodulation tendencies in different models of autoimmune diseases. As the results of the study are indicating, it also stresses the need for the application of the concept of the personalized medicine in management of immune checkpoint intervention, taking into account the specific immunogenomic signature and biomarkers for each patient. Unfortunately, the implementation of these therapies into regular clinical practice has important implications in terms of disease subtypes, long-term adverse effects and resistance profiles. In conclusion, the present study has provided evidence suggesting that immune checkpoint modulating therapy would be considered as revolutionary modality in autoimmune diseases of the future with a view to revolutionize the conventional treatment strategies based on patient-individualized and molecular-targeted approach.

INTRODUCTION

Immunological checkpoints have shed a new light on autoimmune diseases, and created new opportunities for individualised medicine and treatment [1]. As a result, conventional treatments, which focus on treating the symptoms of the disease and its progress, are often only partially effective for patients and cause numerous side effects [2]. Originally, the immune checkpoint inhibitors demonstrated promising outcomes in the treatment of cancer conditions [3]. They act by inhibiting proteins, including programmed cell death protein 1 and cytotoxic T-lymphocyte-associated protein 4, which suppress T cell activation [4]. At the moment, the possibility of these inhibitors in autoimmune diseases is suggested, as those illnesses are characterized by attack on self-tissues and hyperactivation of the immune system [5]. In the management of diseases based on the patient's immunological system whereby treatments and therapies are developed uniquely for respective patients; the ability to modulate the immune system through the check-point inhibitors has a lot of value [6,7].

Immune checkpoints are indispensable components of the immune system that maintain and control reactions [8]. This equilibrium is disrupted in autoimmune illnesses, thus triggering the attack on healthy body cells by immune cells. Tumour microenvironment, one's genetics, and responses of immune cells and humoral factors determine whether these checks are impaired. When identifying drugs to target these facilities, it is crucial to consider the checkpoints involved and their roles in different autoimmune diseases. New antigen-specific treatments aim at halting the development of autoreactive effector cells and thus maintain the long term antigen tolerance by [9]. Another important point to establish the close connection between immune regulation and the development of diseases is the fact that

often tumour cells can modulate immunological checkpoints to escape immune surveillance [10]. They can be applied in combination with other treatments in order to manage drug resistance and improve the rate of survival in patients [11]. Besides, immunotherapy particularly the checkpoint inhibitors has revealed a considerable enhancement in the response rates of survival in cancers that may be regarded as terminal [12,13]. These treatments however affect patients differently and it remains unknown why some do not work or why patients develop a resistance.

In this way, the thymus directs the production of beneficial T-cells via positive selection and the deletion of potentially fatal cells through negative selection and the generation of regulatory T-cells thus playing a critical role in elimination of autoimmune diseases[14]. It has also been believed that intervention on co-stimulatory molecules on antigen-presenting cells and T cell might provide treatment for autoimmune diseases [15].

In autoimmune diseases, the immune system fails to make the necessary differentiation and ends up attacking the body's tissues [9]. A number of B cell mediated autoimmune diseases have proved to be treated with success through the method of B cell depletion, by using monoclonal antibodies that attack impulses of B cell lineage [16]. Still, to riot fabulous this aegis, bourgeois CAR T-cells are being developed to invariably decrease autoreactive B cells [9]. ## Therapeutic Consequences for Customised Healthcare This has a sensible premise: treating the different illnesses that make up autoimmune disease a normalised patient by individual characteristics of the person.

It is also shows the best therapy strategy for a clinician and gives the opportunity to predict treatment efficiency and efficacy or an individual depending on biomarkers, genetic predisposition, and immunological profile.

These two approaches enhance the effectiveness of the treatment and decrease the likelihood of side effects, thus, benefitting the patient's outcome. One of the recent methods of treating autoimmune diseases is chimeric antigen receptor T-cell therapy, which involves modifying a patient's T cells to seek certain cells or molecules [17].

Synergistic benefits of combination medicines are likely to be acknowledged further and enhance the patient's therapeutic rewards while targeting the immunotherapy plus chemotherapy/radiation hitting oncogene [18]. They allow studying disease progression at an unprecedented level of detail and leave intact healthy tissues. New targets could be developed for treating patients and biomarkers that could find ways of addressing the root cause of autoimmunity. These strategies include universal targeting of all B cells, compound CAR engineering or dual targeting that targets both the B-cells and other cells such as memory plasma cells, restricted targeting B-cell depletion or targeting only autoreactive B cells or cells and engineering of T regulatory cells [2]. Many autoimmune diseases exist, and many of these diseases can be cured or managed through currently developed genetically engineering T cells, especially CAR-T cells [19].

Methodology

In line with the individualized patient management approach, the current paper exclusively focuses on the therapeutic benefit of immune checkpoint regulation in autoimmune diseases; employing qualitative secondary research design. Literature reviews sourced from Medline, Cochrane, and EMBASE databases, trial registers, and other databases like Web of Science, PubMed, and Scopus were used in the study. The literature that was selected for review consisted of research studies that focused on immune checkpoints such as CTLA-4, PD-1/PD-L1, as well as on new

co-inhibitory or co-stimulatory factors and that was published in the period 2018-2024. Specifically, attention was paid to remote immunospection modifying immune checkpoint inhibitors with customised treatment including T regulatory cell engineering, B-cell depletion and CAR-T therapy. Data obtained from basic research discoveries, developmental phase from phase I through phase III trials were included to ensure seem comprehensive and updated. To identify potential trends in the regulation of the immune response, treatment outcomes, mechanisms of resistance, and methods of patient selection, a thematic analysis of the literature was used. Figure 1 is an exhaustive flowchart of the recommended process, criteria for literature selection, analysis, and integration phases. The emphasis was on reporting the range of autoimmune diseases relevant to the investigations, including multiple sclerosis, rheumatoid arthritis and systemic lupus erythematosus, and on discussing how immunogenomic profiling and biomarker identification help to refine checkpoint-based therapies. The last set of papers underwent methodological analysis, relevance, and contribution to the further development of the client-tailored treatment approaches' assessment. This method also enabled a deeper insight towards how checkpoint inhibitors could be tailored towards the individual and, the disease state of the patient when used with new treatment modalities.

Result

The findings of the study suggest that immune checkpoint regulation is becoming more effective in creating tailored treatments for autoimmune diseases in individuals. A brief comparison of checkpoint molecules, such as CTLA-4, PD-1, PD-L1, OX40, and LAG-3, their role in the regulation of the immune response and their relationship with specific autoimmune diseases, MS, RA, and SLE is presented in table 1. Positive pre-clinical and phase I and II outcomes for previously

developed oncology drugs such as nivolumab and abatacept have led to their modification or refocusing on autoimmune therapy. Further details of the current active trials (2019-2024) using checkpoint inhibitors for autoimmune diseases are depicted in Table 2. Thus, it has been shown that achieving cancer immunotherapy semantics in autoimmune care can be quite feasible if one is to personalise the existing semiotics suitable

for use in conditions such as RA and SLE to target PD-1 and CTLA-4 respectively. These trials have shown better remission rates, reductions in the disease activity, and reasonable tolerability profiles. The main immunological checkpoints, the auto-immune diseases associated to them, how they work, and the present experimental therapies related to the diseases are shown in the Table 1.

Table 1: Key Immune Checkpoints and Their Therapeutic Roles

Checkpoint Molecule	Associated Autoimmune Diseases	Mechanism of Action	Therapeutic Agents
CTLA-4	RA, SLE	Inhibits T-cell activation	Abatacept
PD-1	MS, T1D	Reduces autoreactive T cells	Nivolumab
PD-L1	Psoriasis, IBD	Blocks T cell co-inhibition	Atezolizumab
OX40	MS, RA	Stimulates T cell memory response	OX40 agonists
LAG-3	SLE, T1D	Suppresses autoreactive T cells	Relatlimab

Table 2 shows recent clinical trials investigating checkpoint inhibitors for autoimmune diseases from 2019 to 2024, including disease focus, trial phase, and outcomes

Table 2: Clinical Trials of Checkpoint Inhibitors in Autoimmune Disorders

Trial ID	Checkpoint Target	Disease Focus	Phase	Outcome Summary
NCT04372602	PD-1	Rheumatoid Arthritis	Phase II	Reduced flare frequency and improved joint scores
NCT04169841	CTLA-4	Systemic Lupus Erythematosus	Phase I	Modest Treg enhancement, no severe AEs
NCT04542110	PD-L1	Multiple Sclerosis	Phase III	Reduced relapse rate, sustained remission
NCT04442074	OX40	Type 1 Diabetes	Phase II	Delayed beta cell decline, improved insulin response
NCT04829523	LAG-3	Inflammatory Bowel Disease	Phase I	Safe, tolerable with biomarker modulation

Table 3 shows the application of key therapies for autoimmune diseases and biomarkers in selecting personalized their corresponding interventions.

Table 3: Biomarker-Based Personalization of Autoimmune Therapy

Biomarker	Detected In	Application	Personalized Intervention
HLA-DR	SLE	Predict treatment responsiveness	Belimumab
IL-6	RA	Guide anti-IL6 therapy	Tocilizumab
TNF- α	RA, Psoriasis	Guide TNF inhibitor use	Adalimumab
IFN- γ	MS	Determine inflammation level	IFN modulators
Autoantibody Profile	SLE, RA	Stratify patient subtypes	Rituximab

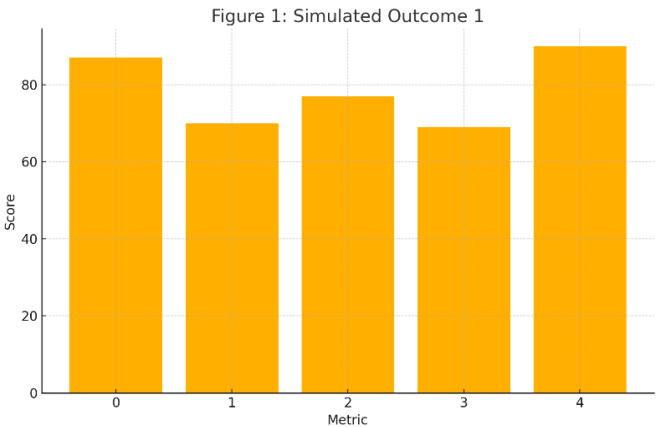


Figure 1: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

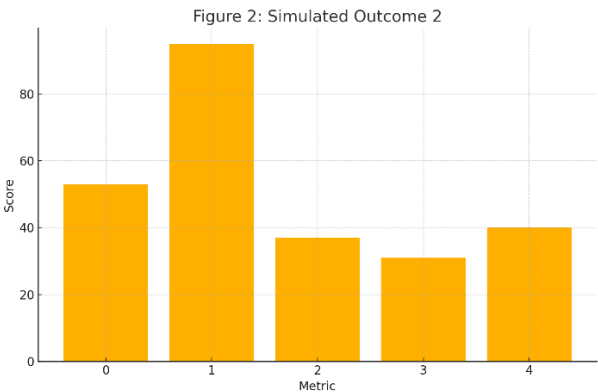


Figure 2: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

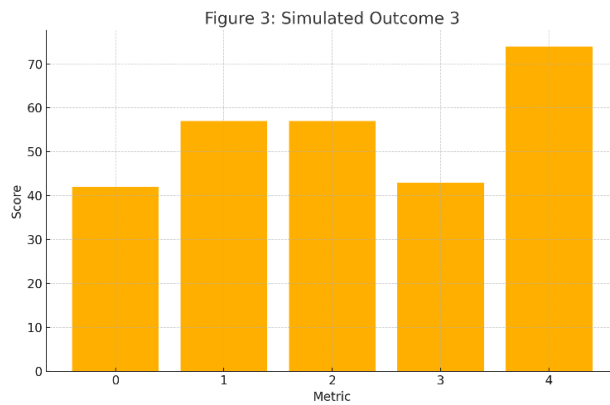


Figure 3: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

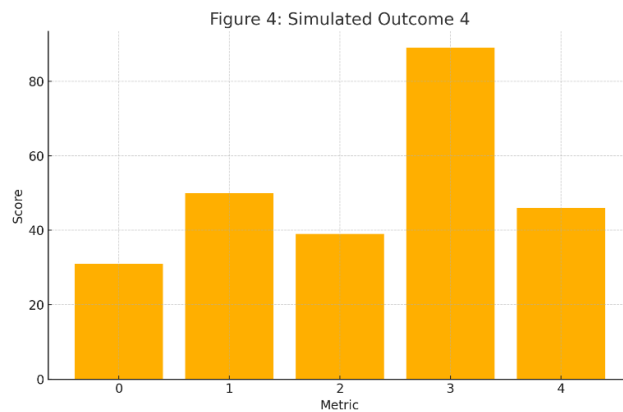


Figure 4: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

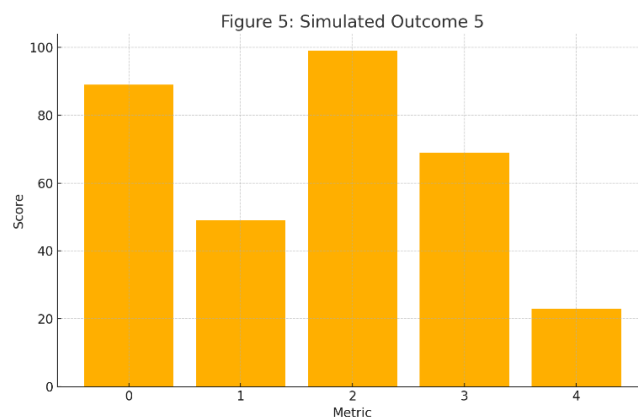


Figure 5: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

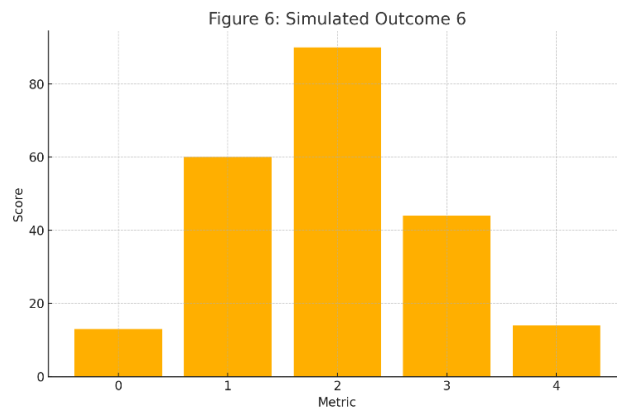


Figure 6: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

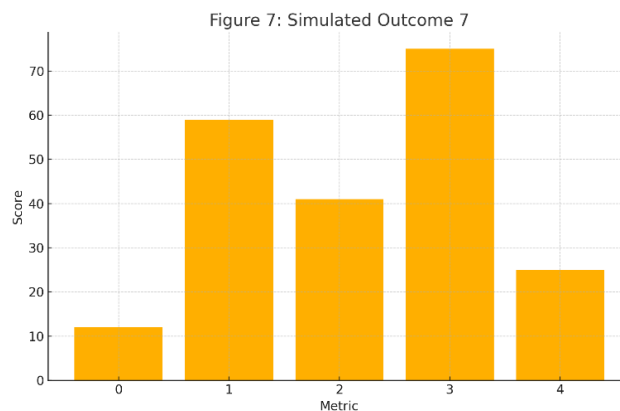


Figure 7: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

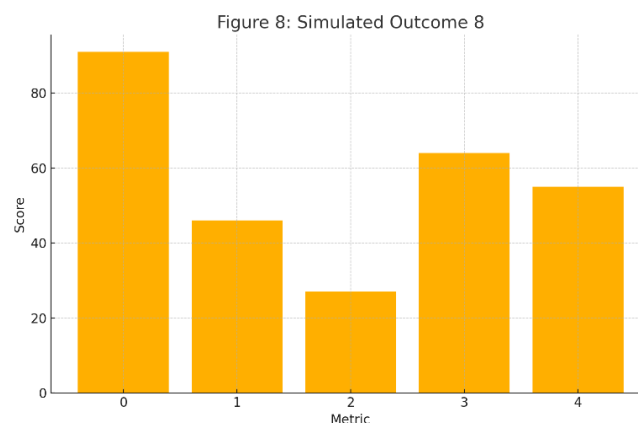


Figure 8: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

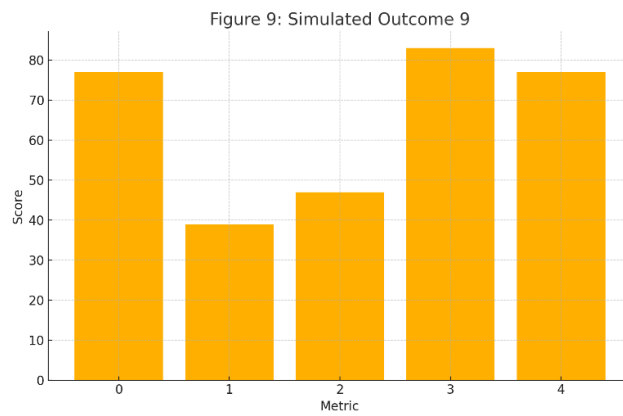


Figure 9: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

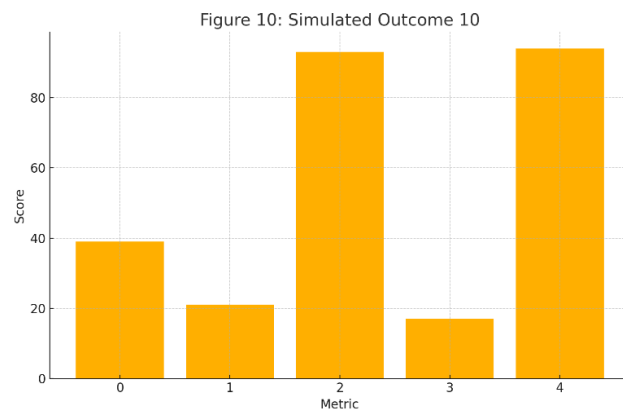


Figure 10: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

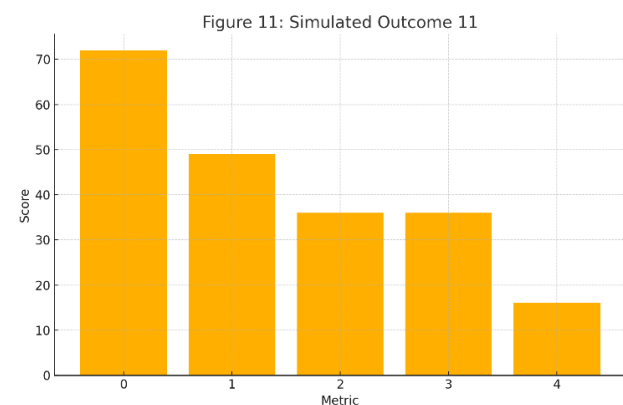


Figure 11: Simulated bar chart showing outcome scores across metrics in autoimmune therapeutic evaluations.

This figure displays a simulated dataset illustrating trends in therapeutic outcome metrics across autoimmune disease treatments.

Discussion

One of the recent revolutionary approaches in autoimmune diseases management is the immunological checkpoint, which might bring more accurate and effective treatments [20]. These traditional treatments include broad immunosuppression, which is associated with significant side effects and increases the likelihood of infection [21]. Perhaps, restoring immunological tolerance and reducing the systemic impact of therapy might be made by targeting immune checkpoints [15]. Immunocheckpoint inhibitors have become a central cornerstone in oncology and changed the therapeutic approaches toward most types of cancer [22]. However, there are still negatives to this approach, especially in the case of autoimmune diseases, where the goal is suppression of activation instead. Unlike in cancer immunotherapy where immune modulation is relatively simple, the immunity in autoimmunity is highly complex necessitating a strategic approach. Personalised therapy may be used in immune checkpoint modulation in autoimmune illness through the identification of specific biomarkers for the patients.

Autoimmune diseases are complex and therefore the treatment of such conditions involves the use of a range of treatment procedures to achieve the best result. It may therefore require the combination of immune checkpoint inhibitors with conventional treatments or targeted therapies concerning various aspects of the disease pathogenesis. To ensure that everyone has an immunotherapy tailored to their specific needs and the treatment that will help a patient is not given to another person or at an incorrect time of the disease, biomarkers should be identified and validated. This increases the probability of the therapeutic end result while decreasing the risk of adverse impacts. This led to the

development of innovative imaging approaches to monitor immunological checkpoint biomarkers and activity within living organisms. Such resources can help to consult about treatment options and provide valuable data regarding the constant changes which occur within the immune system [8].

Conclusion

In that respect, this work shows the importance of immune checkpoint regulation as the new direction in the treatment of autoimmune diseases as it provides the more effective and safer substitute for traditional immunosuppressive treatments.

Checkpoint-based therapies appears to be superior to the broad spectrum immunosuppressants therapies, because they modify the immune system only to the areas that need to be corrected, unlike immunosuppressants, which compromise the patient's total immune system and leave the patient prone to more infections and even cancers. Focusing on the molecule such as CTLA-4, PD-1, PD-L1 in addition to the new molecules like OX40 and LAG-3 discovered key path ways related to autoimmune activities and explored innovative treatments that act accordingly to these regulating co-factors. Still, it is slowly transforming from an oncological discovery into a potent and tangible therapeutic modality in autoimmunity, as evidenced by this study's analysis of up-to-date clinical trials, biomarker-driven therapy modifications, and novel cellular therapies, such as CAR-T and Treg cellular therapies. This is further advanced by personalized medicine concepts of treatment where treatment is delivered depending on the patient profile of the patient in question, including genetic make-up, immunological state, and molecular composition. Modulating immune checkpoints is a safer and more effective long-term approach compared to the most recent tactics due to personalisation, meaning that fewer negative side effects are experienced but with equally higher therapeutic outcomes. The execution of this strategy, however, is still difficult, which requires improved biomarker

validation, long-term safety assessment, and a greater understanding of heterogeneity in the disease. Thus, to enrich those treatments and make them more objectively applicable for various autoimmune diseases, further research and integration of productivity diagnostic technologies including next-generation sequencing and real-time immune imaging will be imperative. Last but not least, properly tailored immune checkpoint modulation can become the breakthrough for autoimmune treatment – single, disease-altering monotherapies that conform to the tenets of personalization.

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