Dendritic cell–derived exosomes for cancer therapy

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DC-derived exosomes (Dex) are nanometer-sized membrane vesicles that are secreted by the sentinel antigen-presenting cells of the immune system: DCs. Like DCs, the molecular composition of Dex includes surface expression of functional MHC-peptide complexes, costimulatory molecules, and other components that interact with immune cells. Dex have the potential to facilitate immune cell–dependent tumor rejection and have distinct advantages over cell-based immunotherapies involving DCs. Accordingly, Dex-based phase I and II clinical trials have been conducted in advanced malignancies, showing the feasibility and safety of the approach, as well as the propensity of these nanovesicles to mediate T and NK cell–based immune responses in patients. This Review will evaluate the interactions of Dex with immune cells, their clinical progress, and the future of Dex immunotherapy for cancer.

Introduction

As the sentinel antigen-presenting cells (APCs) of the immune system, DCs play a central role in initiating antigen-specific immunity and tolerance (1). In cancer, DCs act as the initial link between oncogenesis and the host immune system, the first step of a cancer/immunity cycle that aims to eliminate cancer cells through the activation of T cells (2). Tumor-proximal DCs can capture neoantigens created and released during oncogenesis, which the DCs subsequently process and present to cognate T cells to generate antitumor T cell responses. However, such T cell responses can only be generated if certain additional conditions are met in the local environment (2). These conditions consist of locally present immunogenic signals, such as proinflammatory cytokines, danger-associated molecular patterns (DAMPs), or pathogen-associated molecular patterns (PAMPs). Such signals trigger DCs to present captured tumor-associated antigens (TAAAs) via MHC class I (MHC-I) and MHC-II molecules to T cells in cooperation with costimulatory molecules such as CD80 and CD86, resulting in the priming and activation of TAA-specific effector T cells.

Therapies harnessing these properties of DCs to generate immune responses against tumors have great potential, though clinical progress of this application remains in its infancy. One notable exception is the success of the immunotherapy sipuleucel-T for early-stage, hormone-refractory prostate cancer. Sipuleucel-T is composed of autologous peripheral blood mononuclear cells (PBMCs) including APCs (such as DCs and their precursors) that have been stimulated ex vivo with a fusion protein consisting of the cytokine granulocyte macrophage colony-stimulating factor (GM-CSF), which drives DC differentiation and activation, combined with a prostate antigen (3). Nonetheless, DC-based immunotherapy is challenging to practice in clinical settings. Implementing such therapies across large populations is costly, requires dedicated expertise, and requires monitoring of well-defined quality control parameters. Furthermore, it is difficult to store DCs over long periods of time while maintaining their efficacy (4).

The use of DC-derived exosomes (Dex) has been heralded as a solution to many of the technical challenges associated with DC-based immunotherapy (see Table 1) because they maintain the essential immunostimulatory faculties of DCs (e.g., sharing the ability to present antigens to T cells), while the stable nature of exosomal membranes allows their frozen storage for at least 6 months (5). As biologics, Dex are also more amenable to a strictly regulated and monitored manufacturing process (e.g., their composition and MHC-I and MHC-II content can be easily defined), and they lack the risks associated with viable cellular or viral therapies such as the risk of in vivo replication (6). Finally, treatment with cell-free Dex may be more resistant to immunomodulatory events that occur in tumors than other anticancer vaccines; such events can downregulate costimulatory molecules on DCs and impede stimulation of T cell responses (7).

As discussed in detail in other sections of this review series, DCs are one of the many cell types able to secrete membrane vesicles, such as exosomes, into the extracellular environment. This manner of signaling can modulate recipient cells, such as immune cells or cancer cells, to a level beyond classical ligand/receptor signaling pathways and can create complex cellular modifications that may play a substantial role in how tumor development or immune responses proceed. Moreover, detection of circulating, cancer cell–derived exosomes can serve as a noninvasive diagnostic and screening tool to detect early stages of cancer, facilitating...
deployment of therapeutic interventions (8) (discussed elsewhere in this review series; refs. 9, 10, 11). Secreted membrane vesicles consist of a surrounding lipid bilayer containing various transmembrane proteins; the bilayer encloses various cytosolic components and molecules (e.g., proteins, lipids, and nucleic acids) from the donor cell. While several different subtypes of membrane vesicles have been characterized, such as apoptotic bodies and the microvesicles that directly bud from the plasma membrane (12), the focus of this review will be on the exosomes secreted by DCs, which have progressed to the stage of clinical testing.

The molecular composition of Dex

The molecular composition of Dex allows functionality as an immunotherapy. Similarly to all other exosomes, Dex have a size of 30–100 nm and are initially formed within the cell by the inward budding of endosomal membranes. The resulting endosome formed by this process with its content of intraluminal vesicles is referred to as a multivesicular body (MVB) (13). If MVBs are not targeted for lysosomal degradation, they may fuse with the cell surface membranes of DCs to allow release of their intraluminal vesicles as exosomes (13, 14). The first studies of the Dex proteome revealed a unique molecular composition that allowed for strong immunostimulatory functionality (15, 16). Dex were found to possess MHC-I and MHC-II molecules, which could potentially stimulate CD8+ and CD4+ T cells, respectively, as well as costimulatory molecules (15–18) (see below). Dex possess a variety of surface membrane proteins, including the integrin α and β chains (αMβ2), the immunoglobulin family member ICAM-1, and milk fat globule EGF factor 8 (MFG-E8), which allow for effective targeting and docking to recipient cells (12, 15, 16). MFG-E8 is an abundant protein constituent of Dex membranes that can bind externalized phosphatidylserine on the Dex outer membrane and facilitate uptake of exosomes by linking Dex with αβ3 or αβ5 integrins present on the receiving cell (16). The tetraspanin family of proteins, including CD9, CD63, and CD81, are also well-defined constituents of the Dex surface membrane (15, 16). These and other membrane microdomain organizing proteins are postulated to participate in exosome/acceptor cell interactions (15, 16).

Various cytosolic proteins derived from the donor DC are present on the Dex outer membrane. These proteins appear to be related to Dex biogenesis from DC endocytic compartments (e.g., annexins, RAB proteins, and tumor susceptibility gene 101 [TSG101]) or are participants in signal transduction pathways (e.g., G proteins and kinases). The heat shock cognate protein HSC73 makes up a significant fraction of the cytosolic proteins found in Dex (16). This HSF70 family member, together with members of the HSP90 family that are also present within Dex, may add to Dex immunogenicity, since these proteins can activate various immune cells and have significant roles as antigen chaperones and in MHC loading (18). In addition to these protein families, exosomes also contain mRNA and small RNA (including miR) molecules, which they can transfer between cells (19, 20). RNA transfer via Dex has been shown as a means of communication and posttranscriptional modification between DCs, since exosome-delivered miRs were demonstrated to repress target mRNAs of Dex-accepting DCs (21). These findings could have wide implications for Dex immunotherapy, as they suggest that particular RNA profiles of Dex (or, more specifically, those of the donor DC) could influence Dex immunogenicity through posttranscriptional modifications of acceptor APCs.

Direct and indirect molecular interactions between Dex and T cells. Perhaps the most important attributes of Dex surface membranes, which also set them apart from exosomes secreted by other immune cells, is their possession of DC-originating molecules involved in antigen processing and presentation. It was initially discovered that exosomes derived from B cells possess functional peptide/MHC-II complexes upon their surface membranes and that they could induce antigen-specific MHC-II-restricted T cell responses (22). Dex also have abundant MHC-I molecules (16, 14), which are similarly able to induce T cell responses (23, 24). Importantly, Dex derived from tumor peptide–stimulated DCs were able to prime tumor-specific cytotoxic T lymphocyte (CTL)

### Table 1. Potential functional differences between DC- and Dex-based therapy

<table>
<thead>
<tr>
<th>Practical/functional difference</th>
<th>DC</th>
<th>Dex</th>
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<tbody>
<tr>
<td>Determination of molecular parameters for QC</td>
<td>Molecular composition can change, difficulties defining QC parameters</td>
<td>Dex molecular composition can be strictly defined for each patient (5)</td>
</tr>
<tr>
<td>Abundance of surface peptide–MHC-II complexes</td>
<td>Fewer peptide–MHC-II complexes on the surface membrane, lower yields</td>
<td>Dex are enriched in peptide–MHC-II complexes, by 10–100× that of DCs, allowing up to 6 months Dex therapy from a single leukapheresis (5)</td>
</tr>
<tr>
<td>Stability of vaccine preparation</td>
<td>Living cell, issues with storage and stability over longer periods</td>
<td>Lipid composition of Dex allows high stability for &gt;6 months at ~80°C (5)</td>
</tr>
<tr>
<td>Immunostimulation of NK cells</td>
<td>DCs can fail to express sufficient NK cell receptor ligands (42)</td>
<td>Membrane expression of ligands for NK receptors (42, 43, 45)</td>
</tr>
<tr>
<td>Localization</td>
<td>Rely on chemokine gradients and other signals for access and localization in lymph nodes (54)</td>
<td>Potentially more thorough dispersion within lymph nodes and access to multiple immune cell types (74), not reliant on chemotactic signaling but rather on their surface membrane receptor topography (12, 75, 76)</td>
</tr>
<tr>
<td>Resistance to immunosuppression</td>
<td>Susceptible to immunosuppressive molecules and immunoregulation in the tumor microenvironment (7)</td>
<td>Cannot respond to immunosuppressive molecules</td>
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</table>

QC, quality control.
The constituents of exosome surface membranes, such as integrins and ICAMs, facilitate their binding to and uptake by APCs to enable indirect antigen presentation. Interestingly, the efficiency of indirect T cell stimulation appears to be highly dependent upon the activation status of the donor APC from which exosomes are derived. For example, the exosomes released from DCs treated with LPS or IFN-γ (mature DCs) have increased surface expression of ICAM-1 that presumably enhances their uptake by DCs and increased expression of MHC molecules and CD86, which contribute to T cell activation (31–33). Consistent with their topography, these mature exosomes have a greater capacity to stimulate T cells compared with exosomes derived from immature DCs (24, 31–33).

Following binding to bystander APCs, some of the Dex — but not all individual Dex molecules — are internalized by phagocytosis or macropinocytosis, with the remaining Dex likely retained on the DC surface. The proportion of internalized exosomes is dependent upon the maturation status of the accepting APC. It has been shown that immature DCs internalize exosomes more efficiently than mature DCs, whereas mature DCs are likely to retain more Dex on their surfaces, although presumably surface-retained Dex still interact with T cells (26). Furthermore, priming of naive T cells has been shown to occur only if APCs are present (25, 26, 28).

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peptide complexes can then be transported back to the DC surface for presentation to T cells.

A second mechanism for Dex-mediated indirect antigen presentation to T cells is a process known as cross-dressing where, following binding to APCs, Dex merge with the acceptor APC surface membrane, thereby transferring their peptide/MHC complexes (Figure 1 and refs. 28, 35). This would potentially allow immediate recognition of MHC-presented peptides by T cells without the need for antigen processing and could enable the Dex-transferred antigen presentation machinery to benefit from additional costimulatory molecules and factors present on the accepting APC. In support of this paradigm, Dex can activate T cells only if mature DCs are present, even when mature DCs are deficient for MHC-II (28), suggesting Dex-to-DC cross-dressing of MHC/peptide complexes. Expression of the costimulatory molecules CD80 and CD86 on the presenting DCs was required for T cell activation (28). However, Dex MHC-I cross-dressing of bystander DCs does not appear to induce ovalbumin-specific (OT-I) CD8 + T cell proliferation in an ovalbumin experimental model but instead occurs through Dex internalization and reprocessing of Dex-derived antigen molecules, thereby transferring their peptide/MHC complexes on the acceptor APC surface (Figure 1 and refs. 28, 35). This would potentially allow immediate recognition of MHC-presented peptides by T cells without the need for antigen processing and could enable the Dex-transferred antigen presentation machinery to benefit from additional costimulatory molecules and factors present on the accepting APC. In support of this paradigm, Dex can activate T cells only if mature DCs are present, even when mature DCs are deficient for MHC-II (28), suggesting Dex-to-DC cross-dressing of MHC/peptide complexes. Expression of the costimulatory molecules CD80 and CD86 on the presenting DCs was required for T cell activation (28). However, Dex MHC-I cross-dressing of bystander DCs does not appear to induce ovalbumin-specific (OT-I) CD8 + T cell proliferation in an ovalbumin experimental model but instead occurs through Dex internalization and reprocessing of Dex-derived antigen molecules, thereby transferring their peptide/MHC complexes on the acceptor APC surface (Figure 1 and refs. 28, 35). This would potentially allow immediate recognition of MHC-presented peptides by T cells without the need for antigen processing and could enable the Dex-transferred antigen presentation machinery to benefit from additional costimulatory molecules and factors present on the accepting APC. In support of this paradigm, Dex can activate T cells only if mature DCs are present, even when mature DCs are deficient for MHC-II (28), suggesting Dex-to-DC cross-dressing of MHC/peptide complexes. Expression of the costimulatory molecules CD80 and CD86 on the presenting DCs was required for T cell activation (28). However, Dex MHC-I cross-dressing of bystander DCs does not appear to induce ovalbumin-specific (OT-I) CD8 + T cell proliferation in an ovalbumin experimental model but instead occurs through Dex internalization and reprocessing of Dex-derived antigen molecules, thereby transferring their peptide/MHC complexes on the acceptor APC surface (Figure 1 and refs. 28, 35). This would potentially allow immediate recognition of MHC-presented peptides by T cells without the need for antigen processing and could enable the Dex-transferred antigen presentation machinery to benefit from additional costimulatory molecules and factors present on the accepting APC. In support of this paradigm, Dex can activate T cells only if mature DCs are present, even when mature DCs are deficient for MHC-II (28), suggesting Dex-to-DC cross-dressing of MHC/peptide complexes. Expression of the costimulatory molecules CD80 and CD86 on the presenting DCs was required for T cell activation (28). However, Dex MHC-I cross-dressing of bystander DCs does not appear to induce ovalbumin-specific (OT-I) CD8 + T cell proliferation in an ovalbumin experimental model but instead occurs through Dex internalization and reprocessing of Dex-derived antigen molecules, thereby transferring their peptide/MHC complexes on the acceptor APC surface (Figure 1 and refs. 28, 35). This would potentially allow immediate recognition of MHC-presented peptides by T cells without the need for antigen processing and could enable the Dex-transferred antigen presentation machinery to benefit from additional costimulatory molecules and factors present on the accepting APC. In support of this paradigm, Dex can activate T cells only if mature

Clinical trials of Dex immunotherapies
In view of the high potential and benefits of Dex as an immunotherapy, Dex have been developed for clinical use as cell-free cancer vaccines. Two phase I clinical trials (37, 38) and one phase II trial (39) using Dex have now been completed in advanced cancer patients (see Table 2).

Technical considerations. The manufacturing steps for Dex preparation and purification are detailed in Figure 2. More than 100 individual Dex preparations have been made so far, as the process of vaccine generation is feasible from cells derived from a single leukapheresis. Quality control parameters include the determination of tetraspanins (CD81, CD82, and CD63), high levels of HLA-DR, costimulatory molecules, and exosomal hallmarks such as TSG101 and HSP70 (31). Verification of peptide loading can be performed by incubating Dex with or without HLA-A2 APCs (DCs) and a peptide-specific T cell clone. In this setting, indirect Dex loading of MHC-I-restricted peptides (i.e., loaded onto the donor IFN-γ-treated DCs on day 4 of culture) appears to be as effective as direct peptide loading for stimulation of antigen-specific T cell clones when short peptides were loaded (31). Activation of the latter relative to negative controls confirms functional T cell stimulatory capacity of Dex. For the phase II trial, Dex batch release depended upon a high ratio of tetraspanins and HLA-DR (MHC-II) when compared with control (31, 39). In the phase I trials, Dex MHC-I natural epitopes were acid eluted, whereas no prior elution was performed in the Dex phase II trial. This discontinuation of peptide elution was based on the capacity of high-affinity peptides to compete against and/or coexist with natural epitopes on exosomal surfaces, as shown using ex vivo functional assays with melan-A-specific (MARTI-specific) CTL clones (31, 39).

Phase I trials. The initial phase I trials used Dex produced from autologous monocyte-derived DC cultures loaded with antigenic HLA-presented peptides of melanoma-associated antigen (MAGE). The patients in the two studies received four vaccinations with Dex at weekly intervals.

In the first of these phase I studies (38), nine patients with MAGE-expressing pretreated advanced non–small cell lung cancers (NSCLC) were eligible to receive Dex therapy (Table 2). Three patients who did not exhibit reactivity to MAGE prior to immuni-

Table 2. Dex clinical trials

<table>
<thead>
<tr>
<th>Patient Population</th>
<th>Phase</th>
<th>n</th>
<th>Dex format and approach</th>
<th>TAAs</th>
<th>Effect on immune cell parameters</th>
<th>Trial Outcome</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advanced NSCLC</td>
<td>I</td>
<td>13 (9 completed therapy)</td>
<td>Dex, 4 vaccinations at weekly intervals</td>
<td>MAGE peptides</td>
<td>Limited T cell reactivity. DTH response in 3/9 patients. Possible increase in Treg functions and some NK cell lytic activity.</td>
<td>Safe, well tolerated. Stability of disease in 2 patients with initial disease progression, stability of disease of &gt;12 months in 2/4 initially stable patients.</td>
<td>38</td>
</tr>
<tr>
<td>Metastatic Melanoma</td>
<td>I</td>
<td>15</td>
<td>Dex, 4 vaccinations at weekly intervals</td>
<td>MAGE peptides</td>
<td>No MAGE-specific T cell reactivity. No DTH response. NK cell activation.</td>
<td>Safe, well tolerated. 2 stable diseases, 1 minor response, 1 partial response, 1 mixed response.</td>
<td>37, 42</td>
</tr>
<tr>
<td>Advanced colorectal cancer</td>
<td>I</td>
<td>40</td>
<td>Aex prepared from patient ascites ± GM-CSF, 4 vaccinations at weekly intervals</td>
<td>CEA detected in Aex (inherent loading)</td>
<td>Aex+GM-CS induced CEA-specific CTL responses. DTH induction in both groups.</td>
<td>Safe, well tolerated. 1 stable disease and 1 minor response, both in Aex+GM-CSF group.</td>
<td>40</td>
</tr>
<tr>
<td>Advanced NSCLC</td>
<td>II</td>
<td>26 (22 completed therapy)</td>
<td>IFN-γ-derived Dex, 4 weeks of weekly maintenance immunotherapy after CTX</td>
<td>MAGE-A1, MAGE-A3, NY-ESO, Melan-A/MART1 (all MHC-I–restricted peptides); MAGE-A3, EBV (MHC-II–restricted peptides)</td>
<td>Limited T cell activity. Increased NK cell functions correlating with longer PFS.</td>
<td>7 patients (32%) with stable disease &gt;4 months. 1 patient had a grade-3 hepatotoxicity.</td>
<td>39</td>
</tr>
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EBV, Epstein-Barr virus.
zation showed systemic MAGE-sensitive immune responses one week following the last Dex injection, as determined by delayed-type hypersensitivity (DTH) response. Increases in T cell activity were detected by enzyme-linked immunospot (ELISPOT) assay to MAGE peptides in one of five tested patients. Treg suppression was postulated as a reason for this low rate of T cell activation, since Tregs (defined as CD4+CD25+ T cells) as a percentage of total CD4+ T cells were increased in two of three patients after Dex therapy compared with baseline levels. Interestingly, a potential increase in NK cell lysis ability was also observed in two of four analyzable samples. A good safety profile was the main clinical outcome for the NSCLC phase I study, with disease stability observed in two patients who had disease progression at the start of the study and continued disease stability over 12 months in two of four initially stable patients (38).

The second phase I study enrolled 15 patients with MAGE3+ advanced malignant melanoma (MM), all of whom received Dex (Table 2 and ref. 37). Of these patients, one patient exhibited a partial response to Dex immunotherapy and developed a halo of depigmentation around naevi and disappearance of arterial neovasculature, concomitant with tumor shrinkage. This patient was given continuation therapy with Dex for 4 months, during which time there was stabilization of disease without toxicity. Disease stabilization for up to 24 months was also achieved for another patient who received continued Dex therapy. The overall clinical outcome was two stable diseases, one minor response, one partial response, and one mixed response at skin or lymph node sites, with some of these responses occurring in patients with progressive disease who had previously received other biotherapies or alternative antitumor vaccinations. However, similarly to the other trial, neither MAGE-specific CD4+ and CD8+ T cell responses nor DTH responses could be detected in peripheral blood (37).

A third phase I clinical trial used ascites-derived exosomes (Aex) in combination with GM-CSF to treat 40 advanced colorectal carcinoma (CRC) patients with Aex alone or Aex plus GM-CSF. The patients received four immunizations at weekly intervals (Table 2 and ref. 40). Exosomes were prepared from malignant ascites of the patients and were found to be enriched in MHC-I and MHC-II molecules, costimulatory molecules, and ICAMs. They were also shown to contain the immunogenic carcinoembryonic antigen (CEA) of CRC. A good safety profile was observed for Aex; however, no detectable therapeutic responses were observed except in one patient with stable disease and a second patient who exhibited a minor response after treatment with Aex plus GM-CSF. Interestingly, and in contrast with the low-level stimulation of T cell responses in the Dex clinical studies, Aex plus GM-CSF could induce CEA-specific antitumor CTL responses (40). The majority of the Aex were believed to be derived from CRC cells, although the ascites used in this study likely contained other immune or mesothelial cell-derived nanoparticles or microvesicles. A greater level of TAAs present in Aex compared with Dex may have also been responsible for the greater T cell responses observed in this study as compared with the two Dex phase I trials. However, given findings that tumor-derived exosomes can possess immunosuppressive properties and can facilitate tumor growth, metastasis, and development of drug resistance, care should be taken to distinguish the exact exosome composition of Aex and similar preparations, with a view to removing potentially adverse tumor-derived exosomes when such technologies become available (8, 41).

Interpretation of the immunological findings from the phase I trials: Dex stimulate NK cell functions. The phase I studies confirmed the safety of Dex administration in patients and highlighted the feasibility of large-scale Dex production. Somewhat surprisingly, given the encouraging results from preclinical studies, Dex were
apparently poor stimulators of T cell responses. However, Dex were found to stimulate NK cells.

Following the MM phase I trial, it has been shown that Dex derived from human immature DCs possess killer cell lectin-like receptor subfamily K, member I ligands (NKG2D-L) upon their surface, which can directly engage NKG2D present on NK cells, leading to their activation (42). By using samples from the MM clinical trial, it was found that the number of circulating NK cells significantly increased after 4 weekly vaccinations with Dex. Additionally, NKG2D expression levels and cytotoxicity of NK cells were restored after Dex vaccination in 50% of MM patients presenting with NK cell defects at diagnosis (42). Further investigations of Dex/NK cell interactions in vivo revealed that Dex promote IL-15Ra-dependent NK cell proliferation and an NKG2D-mediated activation of NK cells. These Dex-mediated pro-NK cell effects coincided with improved metastatic control by NK1.1+ cells in C57BL/6 mice inoculated i.v. with B16F10 melanoma cells (42). Dex derived from immature human DCs (and exosomes derived from 293T cells) also express BCL2-associated athanogene 6 (BAG6, also known as BAT3) on their surface (43), which has been identified as a ligand for natural cytotoxicity triggering receptor 3 (NKP30) receptors on NK cells (44). Exosomal BAG6 expression levels were shown to directly correlate with NK cell cytokine release (43). Additionally, Dex surface expression of IFN-γ induces NK cell IFN-γ production (45).

**Phase II trial.** The poor capacity of Dex to stimulate T cell responses in the initial phase I studies encouraged new approaches to improve the interactions of Dex with the adaptive arm of the host immune response. As previously described, the use of exosomes derived from LPS- or IFN-γ-matured DCs (mDex) has been one such new approach, following discoveries that mDex can induce greater T cell stimulation than Dex derived from immature DCs (24, 31, 33). Application of these findings to human DC cultures led to a clinical-grade manufacturing process for Dex vaccines (31) in which IFN-γ stimulation of human DC cultures induces the expression of costimulatory molecules and ICAMs on Dex, yielding a second-generation Dex with enhanced immunostimulatory properties (31, 46). A phase II clinical trial was launched using second-generation Dex in advanced NSCLC patients, with the aim of investigating whether IFN-γ–Dex used as maintenance immunotherapy can ameliorate the rate of progression-free survival (PFS) at 4 months after platinum-based chemotherapy (39).

Twenty-two advanced NSCLC patients who had received four cycles of a first-line platinum-based chemotherapy were eligible to receive IFN-γ–Dex (39). Patients were administered metronomically oral cyclophosphamide (CTX) ahead of IFN-γ–Dex maintenance therapy based upon preclinical (47–49) and clinical evidence (48, 50) demonstrating that this regimen reduces Treg function and stimulates dual IFN-γ/IL-17-producing T cells, thereby facilitating Dex-mediated T cell priming and restoring T cell and NK cell functions. Seven patients (32%) exhibited stable disease after nine injections with IFN-γ–Dex, and these patients continued receiving Dex therapy every three weeks. Consequently, the primary endpoint of the trial, a PFS of 50%, was not reached. No objective response was recorded in the trial, although one patient experienced a long-term stabilization that allowed tumor debulking surgery, a downgrading of disease status by pathologists, and eligibility for local adjuvant thoracic and vertebral radiotherapy.

Regarding the immunological readouts, IFN-γ–Dex immunotherapy again failed to show evidence of TAA-specific T cell immune responses in the patient cohort (39) despite multipetotope loading and the CTX adjuvant. The immunostimulatory capacity of IFN-γ–Dex was instead apparently manifested through augmentation of NKP30-related NK cell functions. Despite low expression levels of NKP30 on stage IV NSCLC patient NK cells, NKP30-stimulated production of IFN-γ and TNF-α by circulating NK cells was increased after four IFN-γ–Dex vaccinations, as compared with the start of the study. Importantly, this increased NKP30-elicited NK cell activation correlated with longer PFS (39).

With regard to a potential mechanism for Dex stimulation of NK cells, it was found that the Dex vaccine preparations possessed the aforementioned membrane-associated NKP30 ligand BAG6, which may have been responsible for activating NK cells of patients in an NKP30-dependent fashion. In support of this hypothesis, BAG6 levels correlated with the MHC-II concentrations of Dex inocula and NKP30-dependent NK cell functions.

This differs from the finding of the previous MM phase I study in which NKG2D (and potentially IL-15/IL-15Rα) signaling was responsible for Dex-mediated NK cell activation (37, 42). Because the Dex immunotherapies utilized in this previous study were not derived from IFN-γ–mDCs in which IFN-γ can upregulate BAG6 expression (51), NKG2D–mediated NK cell stimulation presumably featured more prominently in the absence of BAG6/NKP30 signaling.

**Future directions for Dex immunotherapies**

The early clinical testing of Dex as cancer vaccines has shown limited clinical efficacy in advanced cancer patient cohorts, although cases of disease stabilization were observed in each study. The limited efficacy in the trials might be explained by poor stimulation of adaptive immunity in these patients, potentially due to several factors. A conceivable explanation is the heterogeneity and small size of the patient cohorts, which had received previous anticancer treatments before enrollment.

More importantly, systemic and local immunoregulatory mechanisms (e.g., programmed death ligand-1 [PD-L1] expression in NSCLC cells associated with programmed death-1 [PD-1] upregulation by tumor-infiltrating lymphocytes [TILs], Treg, and myeloid-derived suppressor cell [MDSC] activity) at play in these advanced-stage patients may have also significantly limited this immunotherapy. Another possibility is that the Dex MHC-I– and MHC-II–restricted antigens employed in these advanced patient cohorts were insufficient to drive tumor-targeted T cell responses. The potential of mutated neoepitopes to drive high-avidity T lymphocytes could be exploited for NSCLC patient exosomes (52, 53). Additionally, the in vivo trafficking and fate of Dex in patients is unknown. Instead of reaching the T cell zones of secondary lymphoid organs in sufficient numbers, Dex may be transported to subcapsular sinus macrophages or lymphatic sinus DCs to encounter innate lymphocytes (54). Lastly, it remains to be determined whether Dex express functional PD-L1 or PD-L2 in patients is unknown. Instead of reaching the T cell zones of secondary lymphoid organs in sufficient numbers, Dex may be transported to subcapsular sinus macrophages or lymphatic sinus DCs to encounter innate lymphocytes (54). Lastly, it remains to be determined whether Dex express functional PD-L1 or PD-L2 molecules that may restrict T cell responses.

**Future strategies for development of Dex as an immunotherapy.**

Given the problems encountered within a clinical setting, it will be necessary to develop strategies to enhance Dex activity to create a
successful immunotherapy. Dex can be engineered to enhance particular immunostimulatory characteristics; therefore, it is certainly possible to further improve upon the most recent second-generation Dex. Engineering Dex to have a greater surface expression of various costimulatory molecules, with lower expression of immunoregulatory molecules (e.g., immune checkpoint molecules like PD-L1) may enhance Dex activity. The choice of TAAs adopted in the Dex clinical trials might also be improved upon, perhaps including several different TAAs or mutated neoantigens to combat tumor evolution (2, 53, 55). Increasing the delivered quantity of such TAAs or the addition of GM-CSF to Dex immunotherapies may also help stimulate T cell responses, given the CTL activation observed in the Aex phase I trial (40). Moreover, recent findings have suggested that the design of Dex immunotherapies to stimulate B cell responses alongside T cell stimulation may optimize their immunogenicity (56, 57). Vaccination of mice with protein-loaded Dex, but not T cell peptide–loaded Dex, induced CD8+ T cell responses and control of tumor growth, and these effects depended on the presence of both B cells and CD4+ T cells (56).

Dex immunotherapies may also benefit from the addition of TLR ligand adjuvants. CpG and polyinosinic:polycytidylic acid — which activate TLR9 and TLR3, respectively — induced efficient tumor rejection and Dex-mediated CD8+ T cell priming in a melanoma model using HLA-A2 transgenic mice (58). Thus, the inherent capability of Dex to bind TLR ligands, thereby activating bystander DCs (59), could be implemented to enhance their immunogenicity. In addition to their immunostimulatory capacity, Dex therapies could be engineered to target tumor cells directly (36, 45). Indeed, expression of TNF, FasL, and TRAIL on Dex surface membranes directly triggers caspase activation and apoptosis in tumor cells (45). Engineering Dex to transfer mRNAs encoding relevant neoantigen peptides, or miRs or mRNAs that modulate distinct signaling pathways in immune cells or cancer cells, are other possibilities (20, 21, 60, 61).

Potential next steps for the clinical testing of Dex as an immunotherapy. Dex immunotherapy may be most effective as part of a combinatorial treatment regime. Firstly, combining CTX chemo-therapy with Dex vaccines in vivo significantly enhanced tumor or peptide-induced CD8+ T cell recall responses, leading to a synergistic effect against preestablished tumors (47). However, CTX combination therapy only allowed efficient T cell priming by Dex in humans if genuine adjuvants were present. It is likely that the therapeutic effects seen in this scenario were due to a Dex/CTX therapy retuning of tumor-induced tolerance toward tumor-induced immunogenicity, since CTX can attenuate the suppressive function of Tregs (47, 50, 62, 63). The phase II Dex clinical trial similarly employed CTX to inhibit Tregs prior to Dex immunotherapy (39, 46), although the clinical efficacy of this combination would be more evident in a less advanced cohort. Secondly, combining Dex with PD-1/PD-L1 blockade (or coblockade with anti-CTLA4 therapy) could reduce TIL suppression and enhance T cell priming. Thirdly, a major disadvantage of dealing with NSCLC patients is that the current standard chemotherapeutic regimen does not induce immunogenic cell death (64). Combining Dex with immunogenic cytotoxic drugs would be a suitable option in other malignancies (65, 66). Lastly, it is conceivable that Dex- and NK cell–based immunotherapies (such as anti-killer cell immuno-
globulin-like receptor Ab [anti-KIR Ab]; refs. 67–69) may have synergistic effects against NK cell–dependent malignancies such as gastrointestinal stromal tumors, neuroblastomas, leukemias, and kidney cancers (70–72).

Notwithstanding the many strategies to develop Dex immunotherapy, the predominant reason for the limited efficacy of Dex immunotherapies achieved in clinical trials thus far could be due to the small, advanced cancer-stage patient cohorts used and to the lack of preselection criteria (37, 38). Nonetheless, our phase II Dex trial (39) revealed that patients with detectable serum BAG6 (most probably associated with NKP30 functional defects) are most likely to benefit from Dex therapy. Indeed, patients destined to best respond to Dex appear to be those presenting with defective NK cell receptor expression or functions (notably NKG2D or NKP30), with the predominantly targeted NK receptor dependent upon the specific Dex preparation (e.g., immature versus mature Dex) (39, 42). In prospective clinical studies, these parameters (e.g., killer activated receptors [KARs] and their circulating ligands) should be followed with comprehensive immunomonitoring analyses. Careful patient monitoring for potential immuno-related adverse events is also necessary in future trials involving Dex, despite Dex immunotherapy showing a strong safety profile thus far. This, alongside careful patient screening and selection, will be particularly important for future studies investigating potential combinatorial regimes with Dex (e.g., Dex combined with immune checkpoint inhibitors).

Conclusions

Despite the challenges encountered thus far, the prospect of Dex immunotherapy as a novel cancer treatment remains highly promising. Dex are adept mediators of immune responses and the relative technical ease in manipulating their immunostimulatory characteristics (via the donor DC), along with the advantages over cell-based therapies, ensures their future therapeutic uses. Although Dex immunotherapies produce an encouraging stimulatory effect upon NK cells, which likely contributed to the cases where improved clinical outcome was observed (39), it is clear that improved strategies to overcome the apparent lack of T cell responses are necessary. Focus on this hurdle, as well as on less understood Dex components and functions — such as mRNAs, miRs, transported cytokines, production of lipid mediators (73), and mechanisms through which each of these factors interact with acceptor immune cells — will help Dex immunotherapies to reach their full potential.

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