**Review**

**Immune Responses to Neoadjuvant Chemotherapy in Muscle Invasive Bladder Cancer**

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**Abstract.** The secondary effects of chemotherapy, with bone marrow depression and risk of leukopenia, has traditionally been considered being detrimental for the immune system. However, growing evidence suggests a main role for chemotherapy in antitumor immunomodulation. With reference to cisplatin, which is the basis of neoadjuvant chemotherapy in muscle invasive bladder cancer, four different aspects of immunomodulation has thus far been described; increased MHC class I expression, recruitment and proliferation of effector cells, enhancement of tumor-lytic activity of cytotoxic effectors and downregulation of immunosuppressive actors in the microenvironment. Consequently, the role of chemotherapy in cancer is changing from a therapy solely aimed at inducing tumor cell death, to a potent inducer of immune responses and a potential future major partaker in cancer immunotherapy. This is a great opportunity for the urological community to broaden research in this field in order to increase knowledge, optimize and improve the neoadjuvant regimens of muscle invasive bladder cancer to ultimately improve patient outcome.

Keywords: Urinary bladder neoplasms, cisplatin, neoadjuvant therapy, T-lymphocytes, B-lymphocytes, antigen-presenting cells

**ABBREVIATIONS**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>APC</td>
<td>Antigen presenting cell</td>
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<tr>
<td>CTL</td>
<td>Cytotoxic T cell</td>
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<td>DC</td>
<td>Dendritic cells, functions as APCs</td>
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<td>ICD</td>
<td>Immunogenic cell death</td>
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<td>MHC</td>
<td>Major histocompatibility complex (displays peptide fragments of non-self proteins from within the tumor cell, for the immune system to interact with)</td>
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<td>B cell</td>
<td>White blood cell of the lymphocyte subtype. Secrete antigen</td>
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MDSC  myeloid-derived suppressor cells which possess strong immunosuppressive regulatory functions

Tcell  white blood cell of the lymphocyte subtype. Major player in the cell-mediated immunity. Different subsets with different functions.

TCR  T cell receptor

T reg  T regulatory cells, which are considered to maintain strong immunosuppressive regulatory functions

NEOADJUVANT CHEMOTHERAPY IN MUSCLE INVASIVE BLADDER CANCER

The proposed rationale for neoadjuvant chemotherapy in muscle invasive urinary bladder cancer (MIBC), is the early treatment of undetected micro-metastatic disease prior to major surgery for maximum efficacy [1]. Neoadjuvant cisplatin based combination chemotherapy (NAC) has been proposed to be of value for downstaging of the primary tumor and facilitating surgical resection during radical cystectomy (RC), however, the overall survival benefit is only 5–8% in intention-to-treat settings [1–3]. In contrast, complete responders (pT0N0M0) gain substantial survival benefits from NAC [4, 5], and the early identification of responders versus non-responders to NAC therapy is therefore an important area of research. Furthermore, retrospective evaluations of larger populations have revealed that NAC therapy is a positive prognostic factor also with regard to pTis, pTa and pT1 outcomes, similar to the findings in patients with pT0N0 [6]. Interestingly, a growing body of data suggests that NAC efficacy is not only due the induction of tumor cell death following the treatment, but also an anti-tumoral induction of the immune system. However, a solid connection between significant tumor downstaging and NAC-induced/enhanced immune responses, has yet to be demonstrated.

TUMOR IMMUNOLOGY

Mutations are at the center in the multistep process of cancer development, resulting in the formation of foreign proteins with the potential to be recognized by the immune system. The massive expansion of the tumor immunology field during the past decade, is illustrated by the update of the Hallmarks of Cancer theorem, to include tumor immune evasion as an emerging Hallmark [7]. Cancer immune surveillance [8] is the idea that the immune system continuously monitors and protects the body from developing cancer cells. In a proposed process called immunoediting, a malignancy undergoes three phases: (1) The elimination phase, during which immune surveillance prevails. (2) The equilibrium phase, where interactions between tumor cells and the immune system result in evolutionary pressure and “editing” of the tumor phenotype, and (3) the escape phase during which the tumor has acquired properties (i.e. escape mechanisms) allowing it to evade the immune system and grow [8, 9].

Tumor escape mechanisms

Tumor evasion of anti-tumor immune responses constitutes a challenge for future therapeutic strategies. Indeed, many of the latest drug introductions in cancer therapy directly target the natural breaks of the immune system, the most prominent examples being check-point inhibitors such as the monoclonal anti-CTLA4 antibody ipilimumab and anti-PD1/PD-L1 therapies. A wide variety of tumor escape mechanisms have been described including altered antigen presentation, production of immunosuppressive factors and recruitment/induction of regulatory immune cell populations. Here follows a brief introduction to some of the most well studied tumor immune escape mechanisms.

Downregulation of MHC class I

MHC class I (MHCI) complexes present peptides from inside the cell on the cell surface enabling the recognition of altered antigens (so called neoantigens) by cytotoxic T cells (CTLs). Activation of CTLs triggers the release of effector substances such as granzyme and perforin, which in turn eliminate the tumor cells. To survive, many tumors down-regulate or alter their MHCI expression through a variety of different mechanisms such as deletion of the β2-microglobulin genes, direct suppression of MHCI gene transcription or interference with the antigen presentation machinery such as the transporter associated with antigen processing (TAP) [10, 11].

Induction of check point inhibitors

Feedback loops are built into a normal immune response, in order to regulate the length and the mag-
Fig. 1. T cell activation. Schematic illustration of the three signals required to activate a naïve T cell: 1) TCR stimulation by recognition of peptide-MHC complex. 2) Costimulation through e.g. CD28 interaction with B7 molecules such as CD80 and CD86. 3) Cytokines that influence the type of immune response developed.

magnitude of immune activation. Once a naïve T cell recognizes a tumor derived peptide presented by an MHC molecule in a tumor draining lymph node, additional costimulatory signals such as CD80 and CD86 are needed for T cell activation (Fig. 1). During the progression of the immune response, activation is followed by upregulation of inhibitory receptors, such as CTLA-4 or PD-1 that compete with the T cell activation signals, leading to a diminished T cell response. Of note, tumors may induce and enhance the expression of CTLA-4, and PD-1/PD-L1, thus suppressing immune activation [12, 13]. The importance of these immune check points has been demonstrated during recent years by the development and clinical introduction of check point inhibitors such as ipilimumab and nivolumab, targeting these specific pathways [14, 15].

The tumor microenvironment

The tumor microenvironment is where the interactions between cancer cells and the immune system occur and it is very much connected to different modes of tumor immune escape. For example, tumors can actively express immune suppressive factors such as the enzyme indoleamine 2, 3-deoxygenase (IDO) [16], secrete immune modulating cytokines such as TGF-β and IL-10 [11, 17], and moreover the relatively hypoxic environment within the tumor, can constitute an immunosuppressive factor in itself [18].

CYTOTOXIC CHEMOTHERAPIES AND THE IMMUNE SYSTEM

Cancer chemotherapy has traditionally been considered detrimental to the immune system, and had until recently not been evaluated in a broader context. Chemotherapy associated bone marrow depression with ensuing lymphopenia has been interpreted as a hallmark of a weakened immune system with ensuing risk for negative effects on long time survival. However, evidence suggests that lymphopenia may create prime conditions for restarting or re-booting large parts of the immune system through so called homeostasis-driven proliferation of immune cells (promoted by cytokines such as IL-7 and IL-15). The end effect being a diminished tolerance for the altered self-characteristics of tumor cells [19, 20].

GENERAL ASPECTS ON CHEMOTHERAPY-INDUCED ANTITUMOR IMMUNOMODULATION

Chemotherapy and immunogenic cell death

Interestingly, chemotherapeutic drugs have been shown to induce so called immunogenic cell death (ICD) in tumor cells, ultimately resulting in augmented immune responses in tumor vaccination experiments [21]. The tumor cell goes through different phases during the process of ICD, starting from the induced pre-apoptosis over apoptosis, to the final necrotic cell. In all three phases, the immune system interacts with damaged tumor cells in a concerted effort to eliminate the tumor. During the first phase, the DCs of the host, attempt to engulf the pre-apoptotic cell. This due to the tumor cell, after chemotherapy induced damage, starts to expose calreticulin (CRT) on the cell surface. CRT exposure seems to determine the engulfment of dying tumor cells by DCs, and can serve as an DC-activation signal [22]. In the apoptotic phase, the tumor cell releases ATP which attracts more DCs and further leads to a process of DC maturation. In the final necrotic phase, the cell membrane becomes more permeable, leading to release of HMBG-1, leading to further triggering of DCs and to enhanced T cell immunity activation [23, 24]. Interestingly, different chemotherapeutic regimes appear to vary in their ability to induce ICD, suggesting that optimization of clinical chemotherapy protocols could lead to more ICD of tumor cells in patients and consequently better immune activation and clinical response to treatment [21].

CISPLATIN IN THE CONTEXT OF IMMUNOMODULATION

Since cisplatin is the main player in chemotherapy of MIBC, the following sections will focus on the
immunomodulatory effects described for this drug in different settings. Of note, cisplatin as monotherapy in MIBC is considered less efficient than combination therapy, the latter being the Gold standard following the results of the randomized prospective trials [2, 3] and also in a major meta-analysis of all RCTs addressing the concept [25]. Apart from the listed immunomodulatory effects of cisplatin, other components in MVAC as well as in gemcitabine stand-alone, have also been shown to render antitumorous immunological effects [26, 27].

As of today, much work still remains to be done regarding the specific immunomodulatory effects of chemotherapy in MIBC.

Basic immunomodulatory effects of cisplatin

1. *Increased MHC class I expression*
   
   Different research groups have demonstrated that cisplatin enhances cell expression of MHC I in experimental studies of tumor cell lines [28–31]. As discussed previously, MHC I is vital for tumor cell recognition and elimination by CD8\(^+\) cytotoxic T cells, and increased expression would thus accordingly promote anti-tumor immune responses.

2. *Recruitment and proliferation of effector cells*
   
   Homing of effector cells to the actual target area (i.e. to the primary tumor or metastatic deposits) is of primary importance for an effective immune response to occur. Experimental investigations indicate cisplatin as a facilitator and promoter in the homing process [32, 33].

   Interestingly, a combination of cisplatin and gemcitabine in a murine model, investigating mesothelioma and lung cancer tumors, led to substantially increased amounts of activated CTLs, both systematically and locally in the tumor [34].

3. *Enhancement of tumor-lytic activity of cytotoxic effectors*
   
   Pretreatment of CTLs with cisplatin has been shown to increase CTL-mediated killing of tumor cells in lymphoblastoid cell lines [35]. Furthermore, it seems cisplatin can sensitize tumor cells to CTL-mediated killing as demonstrated in a colon carcinoma cell line, in which the mechanism was mediated through the upregulation of both ICAM-1 and Fas expression leading to an increase in caspase 3-like protease activity and apoptosis [36]. Moreover, in a study of human lung carcinoma cell lines with sublethal exposure to cisplatin/vinorelbine, the sensitivity to perforin/granzyme mediated CTL-killing was also significantly increased [37]. Another study of cisplatin in combination with adoptive T cell immunotherapy illustrated that cisplatin-treated tumor cells had increased perforin-independent permeability, leading to increased CTL-mediated Granzyme B activity, and subsequent lethal effects on the tumor cells [38].

4. *Downregulation of immunosuppressive actors in the microenvironment*
   
   Cisplatin has been shown to downregulate both myeloid-derived suppressor cells (MDSC) and T regulatory cells (Treg) in separate studies, with postulated secondary effects on immunocompetent cells being unhampered in their respective activity [29, 32, 37]. Lesterhuis and associates have shown that exposure to platinum based chemotherapy markedly reduced PD-L2 expression (T cell inhibitory molecule programmed death receptor ligand 2), and the effect was determined both on DCs as well as tumor cells. The PD-L2 downregulation resulted in enhanced antigen-specific proliferation and Th1 cytokine secretion, which further improved recognition of tumor cells by T cells [39].

   Experimental data indicates that cisplatin also has a function of displaying and presenting subdominant epitopes of tumor antigens. Thus, enhancing the possibility of the adaptive immune system (CTLs) to recognize otherwise hidden tumor targets. This is especially important due to the risk of immune evasion of an effective immune response towards the dominant epitope [40].

CHEMOTHERAPY-INDUCED ANTI-TUMOR RESPONSES IN MIBC

As mentioned previously, activation of T cells needs two signals, one provided by the MHC-tumor peptide complex and one by costimulatory molecules such as B7.1 (CD80) and B7.2 (CD86). In addition, a third signal mediated by cytokines is responsible for optimal T cell activation. Our group previously demonstrated that cisplatin induces release of the T cell activating cytokine IFN-\(\beta\) by APCs enhancing T cell activation [41]. Recently we also demonstrated that doxorubicin treatment of B cells results in increased expression of the costimulatory molecule CD86 and increased T cell activation [42]. Blocking CD86 with a monoclonal antibody inhibited this T cell activation. In addi-
tion, doxorubicin resulted in decreased production of the anti-inflammatory cytokine IL-10. To investigate the immunological response to doxorubicin, MIBC patients treated with NAC were investigated, demonstrating an increased expression of the costimulatory molecule CD86 of B cells. Thus, doxorubicin given as part of combination-NAC seems to support activation of T cells and anti-tumoral immune responses [42].

FUTURE OPTIONS AND DIRECTIONS

Although tumors can be defined by patient characteristics, histology and staging, this is not sufficient to fully understand an individual’s treatment options, especially not with regards to upcoming immunotherapy approaches. It is important to account for the clinical and immune characteristics of the tumor as well as the patient’s immunological status, not only on a group level, but ultimately for each individual patient. New evolving bladder cancer taxonomies from different research groups [43–45] need to be related to corresponding immunological properties, both in the different subgroups as well as in relation to both innate and adaptive immune responses.

The different compartments in which the tumor cells interact with the immunological defenses also warrant further investigation that includes, apart from the primary tumor, also blood, urine, bone marrow and tumor draining lymph nodes [46, 47].

The immunomodulatory effects of chemotherapy in bladder cancer that have been suggested during recent years, suggest that further research is needed to evaluate the best strategy for optimal administration of chemotherapy and its relative timing to the ensuing RC. A tailored approach would require individual optimization and we propose that there might be major advantages of monitoring each patient’s immune responses regularly during the course of NAC-administration. Hypothetically, an inflexible general timetable with strictly predefined doses may otherwise lead to a suboptimal outcome in terms of endpoints related to immune induction, efficacy on micro-metastatic disease and even long term survival. The starting point should be immunological characterization already at time of primary diagnosis, pre- and post-TURb, in between NAC-cycles, pre- and post-RC and during follow up. Monitoring immunological status, developing and evaluating biomarkers of immunogenic cell death [48] opens up for individually adapted chemotherapy regimens to optimize the stimulatory effect of chemotherapy on the immune system.

CONCLUSIONS

Chemotherapy, once considered only an inducer of tumor cell death, can now be investigated, measured and utilized as possible tool for cancer immunotherapy. There are great opportunities for the scientific urological community to broaden the research in this field, in order to improve our knowledge and optimize neoadjuvant regimens in the field of muscle invasive bladder cancer.

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CONFLICT OF INTEREST

The authors have no conflicts of interest to report.

REFERENCES


