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**Prognostic significance of tumor immune microenvironment and immunotherapy: novel insights and future perspectives in gastric cancer**

Lazăr DC *et al.* Immunotherapy in gastric cancer

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**Abstract**

Despite a decrease in gastric cancer incidence, the development of novel biologic agents and combined therapeutic strategies, the prognosis of gastric cancer remains poor. Recently, the introduction of modern immunotherapy, especially using immune checkpoints inhibitors, led to an improved prognosis in many cancers. The use of immunotherapy was also associated with manageable adverse event profiles and promising results in the treatment of patients with gastric cancer, especially in heavily pretreated patients. These data have led to an accelerated approval of some checkpoints inhibitors in this setting. Understanding the complex relationship between the host immune microenvironment and tumor and the immune escape phenomenon leading to cancer occurrence and progression will subsequently lead to the identification of prognostic immune markers. Furthermore, this understanding will result in the discovery of both new mechanisms for blocking tumor immunosuppressive signals and pathways to stimulate the local immune response by targeting and modulating different subsets of immune cells. Due to the molecular heterogeneity of gastric cancers associated with different clinico-biologic parameters, immune markers expression and prognosis, novel immunotherapy algorithms should be personalized and addressed to selected subsets of gastric tumors, which have been proven to elicit the best clinical responses. Future perspectives in the treatment of gastric cancer include tailored dual immunotherapies or a combination of immunotherapy with other targeted agents with synergistic antitumor effects.

**Key words:** immunotherapy; immune checkpoints inhibitors; tumor immune microenvironment; prognostic significance; gastric cancer

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**Core tip:** The use of modern immunotherapy, including adoptive cells therapy, vaccines, and especially immune therapy using checkpoints inhibitors, has led to encouraging results in clinical trials including gastric cancer patients. This review analyzes the relationship between immune microenvironment profile of the host and tumor development, identification of the immune prognostic markers and future perspectives of immunotherapeutic strategies. The treatment algorithm should be adapted to the specific molecular profile of the gastric cancer subtype in order to obtain maximum clinical benefits.

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**Introduction**

Early estimates situated gastric cancer in the first place, as being the most frequent neoplasia (1975). Despite a decrease in gastric cancer incidents during the last decades, it remains a major health problem globally, with almost one million new cases diagnosed in 2012 (6.8% of the total), after tumors of the lung, breast, colorectum and prostate. There is a high geographical variation in gastric cancer incidence. Approximately 70% of cases occur in developing countries; half of the total number occurs in Eastern Asia, especially China; the incidence rates are approximately twice in men *vs.* women. Overall, this type of tumor represents the third leading cause of cancer death in both sexes, accounting for 723000 deaths in 2012 (8.8% of the total number of cases). The highest mortality rates are seen in Eastern Asia, whereas the lowest rates occur in Northern America; also, high mortality rates are encountered in Central and Eastern Europe and in Central and South America, respectively[1].

Most gastric cancers are diagnosed at an advanced stage, whereas another 25%-50% of cases will develop metastases during the outcome of the disease. Although surgical resection remains the main treatment with curative-intent in gastric cancer patients, there is a poor associated 5-year survival rate, of approximately 20%-25%. Therefore, additional treatments (neoadjuvant/adjuvant), such as chemotherapy and radiotherapy where associated with tumor resection, unfortunately lead to only modest survival benefits. In advanced stages, approximately 50% of cases present local/systemic recurrence after adjuvant treatment, and only 10%-15% of cases achieve a 5-year overall survival[2]. In the metastatic stage, the backbone of treatment is represented by palliative chemotherapy, associated with a poor median overall survival, of approximately 8-10 mo **[**3]. Despite recent advances using novel biologic therapeutic agents, with the exception of trastuzumab (anti-HER2 monoclonal antibody) and ramucirumab (fully humanized monoclonal antibody receptor antagonist to bind VEGFR-2), showing beneficial results by improving overall survival (OS), and therefore approved in first-line (in association with standard chemotherapeutic regimens) and second-line settings, respectively (as monotherapy, or in association with chemotherapy), in advanced and metastatic gastric cancers, clinical trials assessing other targeted agents showed disappointing results in gastric cancer[4-6].

Recently, the therapeutic algorithm and prognosis of many tumors changed radically by introducing immunotherapy, especially using immune checkpoints inhibitors, and the first drug of this class approved by the United Stetes Food and Drug Administration (FDA) was ipilimumab, an anticytotoxic T lymphocyte antigen-4 (CTLA-4) antibody, used in the treatment of advanced melanoma (2011)[7,8]. Afterwards, immune checkpoints inhibitors, which are antagonists of the programmed death (PD)-1/PD-ligand 1 (PD-L1) pathway, were approved by the FDA for the treatment of different tumors, such as melanoma, non-small cell lung cancer (NSCLC), urothelial/renal cell carcinoma, squamous cell carcinoma of the head and neck, Merkel cell carcinoma and Hodgkin’s lymphoma[9].

**Molecular classification of gastric cancer**

The following main histological classifications of gastric cancer have routinely been used: the World Health Organization (WHO) classification[10] that categorizes four histological subtypes, namely, papillary, tubular, mucinous and poorly cohesive, and Lauren`s classification, dividing gastric cancers into intestinal, diffuse and mixed type[11].

Because these two classifications are not able to direct specific therapeutic strategies and additionally, because the group of gastric cancers includes heterogeneity of tumors, there was a need to elaborate new classifications capable of stratifying patients regarding tumor behavior, prognosis and response to specific treatments. For the first time, the molecular assessment of gastric cancer patients was proven to add benefits in the context of the TOGA trial in which a combined treatment with classical chemotherapy and trastuzumab showed an improvement of survival in the subgroup of patients overexpressing HER2[4]. Moreover, the behavior of the tumor and the outcome proved to be different in cases of Asian patients *vs* Caucasians included in several clinical trials[12].

In 2013, Singapore researchers identified three different molecular subtypes of gastric cancer: proliferative (high genomic instability, TP53 mutation), metabolic (high response to 5-FU chemotherapy), and mesenchymal (stem cell-like cancers that are sensitive to PIK3CA-mTOR inhibitors)[13].

The aim of “The Cancer Genome Atlas (TCGA)” project (2014) was to develop a new molecular classification of gastric cancer with clinical impact and to identify the main dysregulated pathways of each subtype of gastric tumors. The TGCA research group divided gastric cancer into four genomic subtypes:

**Chromosomal instability (CIN):** includes approximately 50% of cases, most of the tumors are located at the gastro-esophageal junction or cardia[14]; leads to a loss or gain of some oncogenes and tumor suppressor genes[15]; has a high frequency of TP53 gene and receptor tyrosine kinases mutations and amplifications of cell cycle genes[16]; and has amplifications in oncogene pathways (HER2, BRAF, EGFR, MET, and RAS)[17].

**Microsatellite instability (MSI):** tumor testing methodologies include immunohistochemistry for abnormal absence of MMR protein expression or polymerase chain reaction (PCR) for MSI analysis, to evaluate for MSI-H on a tumor specimen. Immunohistochemistry can predict which gene is most likely to be mutated, the gene for the affected protein. Interpretation of immunohistochemical reports can sometimes be confusing, as “positive” should mean the abnormal absence of MMR protein expression, in contrast to normal presence of expression. MSI testing panels may consist in mononucleotide and dincleotid markers. For classifying MSI, a panel of five markers for the analysis of MSI was recommended by the National Cancer Institute, including two mononucleotide and two dinucleotide repeats. In case that ≥ 2 of the five markers show instability, the genotypes are grouped into high‑frequency (MSI‑H); when only one marker shows instability, into low‑frequency (MSI‑L), and when no marker shows instability, into micro­satellite stable (MSS). MSI‑H consists in 30%‑40% instability markers, while MSI‑L in < 30%‑40%. Bethesda Guidelines have stated that MSI‑H can be defined when instability is present at a mononucleotide loci and MSI‑L when instability is limited at only a dinucleotide loci; mononucleotide repeats were demonstrated to be more sensitive *vs.* dinucleotide loci in detecting MSI. Some studies have shown that both immunohistochemistry and MSI are cost-effective and useful for selecting high-risk patients. A review showed that the sensitivities of MSI and immunohistochemical testing are 77% to 89%, and 83%, respectively; specificities are 90% and 89%, respectively. Some patients may have MSI or abnormal immunohistochemistry due to sporadic development of cancer, rather than an underlying genetic (germline) mutation. MSI accounts for 15-30% of gastric cancers; most often includes tumors of intestinal type, antral location, females and older patients[18,19]; shows mutations in DNA mismatch repair genes, such as MLH1 or MLH2, that lead to the dysfunction of DNA mismatch repair enzymes[20]; is reported in a meta-analysis that demonstrated a 37% reduced mortality risk and prolonged OS in gastric cancer patients with MSI-high (MSI-H) *vs* MSI-low (MSI-L)[21]; is associated in non-colorectal cancer with an increased frequency of somatic mutation and amplification of tumor antigens; therefore, an increased sensitivity to treatment with PD-1 immune checkpoint inhibitors[22,23]; has increased intratumor infiltrate[24]; is reported in studies that show an increased activity of pembrolizumab in gastric cancer[25] and assess the efficacy of nivolumab ± ipilimumab in MSI-H gastrointestinal cancers[26]; and is possibly being considered in the development of a preventive vaccine using neopeptides affecting MSI carcinogenesis[27].

**Genomic stability:** accounts for approximately 20% of gastric cancer cases; has an increased frequency of the diffuse type; indicates main somatic genomic alterations: CDH1, ARIDIA, RHOA; and presents a recurrent interchromosomal translocation involved in cellular motility[28].

**Epstein Barr virus (EBV)-associated:** EBV is detected by *in situ* hybridization or PCR in approximately 10% of gastric cancer patients[29]; it seems to increase 10 times the gastric cancer risk, especially in Far East Asian patients[30] and is more frequent in younger patients[29]; it has a better response to immunotherapy and better survival[31]; it shows that the PD-L1 gene is frequently amplified; approximately 15% of EBV(+) tumors present amplification of chromosomal region 9p24.1 (locus of PD-L1, PD-L2)[32]; EBV (+) is found in approximately 50% of tumor cells and 94% of immune cells; therefore, PD-1 testing could predict a response to immunotherapy in this subset of patients[12]; and it shows PIK3CA mutations that could be targeted using PI(3)-kinase inhibition, DNA hypermethylation and JAK2 mutations[28].

Taking into account specific characteristics of the four subtypes of gastric cancer, it was highlighted that EBV-associated and MSI categories are associated with the best responses to immune therapeutic strategies.

The Asian Cancer Research Group (ACRG) proposed a molecular classification of four molecular subtypes for gastric cancer (2015): one subtype was related to the epithelial-to-mesenchymal transition (MSS/EMT) phenotype, which was associated with highest recurrence frequency, and another subtype was related to the phenotype of microsatellite instability (MSI), cytokine signaling, cell proliferation and methylation signals, including hyper mutated tumors, which was associated with the best overall prognosis. The remaining of the non-EMT and non-MSI patients, were further divided into microsatellite stability (MSS)/p53− and MSS/p53+ molecular subtypes and were associated with an intermediate overall prognosis and recurrence[33,34].

Comparing TCGA *vs* ACRG subtypes of gastric cancers, one may notice a resemblance between MSI tumors, GS and MSS/EMT subtypes, EBV and MSS/TP53+ subtypes, and CIN and MSS/TP53- tumors, respectively[13].

The development of genotyping different subtypes of gastric cancer will provide a guide to molecular targeted drugs that should be investigated in large clinical trials on specific subsets of gastric tumor patients in the future.

**Host immune response in gastric cancer patients**

***Antitumor immune response of the host***

The term “immunosurveillance” refers to the capacity of the host immune system to identify as “non self” the tumor cells, and subsequently to kill them[35]. The immune response includes both innate immunity (represented by macrophages, dendritic cells, and natural killer cells), and specific adaptive immunity (T and B lymphocytes).

The host protective response and the capacity of the tumor to surpass the host immune response are defined as “cancer immunoediting”, which is a process comprising three progressive steps:

**elimination phase:** in this stage, natural killer (NK) cells and T lymphocytes (helper and cytotoxic) secrete interferon IFNγ, leading to a reduction of angiogenesis and proliferation of cancerous cells; moreover, macrophages and dendritic cells (DC) secrete cytokines that activate immune cells to phagocytize dead tumor cells

**equilibrium phase:** residual cancerous cells remain in a dormancy state because DC and cytotoxic T cells secrete IFNγ and inhibitory cytokines (IL12), suppressing them.

**escape phase:** tumor cells change their features which will be transmitted to the daughter cells, therefore escaping immunosuppression and proliferating, along with the apoptosis of the effector immunocytes[36].

This process is illustrated in Figure 1.

The tumor mechanisms to evade suppression by the immune system may include a reduced expression of the tumor antigens and major histocompatibility class I (MHC) antigen, Fas-L modulation, increased synthesis of inhibitory cytokines such as TGFß1, IL10, IL6, VEGF, prostaglandin, and an increased number of T regulatory lymphocytes (Treg) and myeloid-derived suppressor cells (MDSC)[37].

***Tumor immunosuppressive microenvironment***

The most important components of the tumor immunosuppressive microenvironment are represented by the regulatory T cells (Tregs) and mesenchymal- or bone-marrow-derived stem cells (BM-MSCs). *T regs* represent CD4+CD25+FOXP3+ T lymphocytes that determine reduced activity of cytotoxic and helper T cells, and of NK cells and are involved in the immunological tolerance to self-antigen and the persistence of H. pylori-related inflammation[38]. *BM-MSCs* migrate to cancerous tissues, and, in some animal models, were shown to create an immunosuppressive environment in chronic H. pylori infection and to represent a “seeding point” for gastric carcinogenesis[39-41].

In this regard, immunotherapeutic strategies directed against Tregs and BM-MSCs and against immunosuppressive cytokines seem to be promising.

Cytotoxic T-lymphocyte-associated antigen 4 (CTLA-4) and programmed death-1 (PD-1) play essential roles in the immune checkpoint modulation. Normally, these molecules modulate the T-lymphocytes response to antigens. CTLA-4 represents an inhibitory receptor exhibited by T cells, whereas PD-1 represents a co-inhibitory receptor located on the cell surface, suppressing T-cell activity in peripheral tissues in the context of inflammation. PD-1 is widely expressed on T and B lymphocytes, monocytes, and natural killer cells; conversely to CTLA-4, PD-1 is involved in subsequent phases of immune responses[42]. It has been shown that various tumor cells upregulate PD-L1. In gastrointestinal cancers, PD-L1 upregulation has been identified in pancreatic, gastric, and colorectal cancers, correlating in several studies with a poor prognosis[43,44].

***Significance and prognostic role of tumor-infiltrating lymphocytes (TILs) in gastric cancer***

TILs comprise the presence of T cells, B cells and NK cells[45]. T cells include cytotoxic lymphocytes CD8+, helper T cells CD4+, memory T cells CD45 RO+ and T regulatory cells FOXP3+. Specific cell membrane antigens of TILs bind to specific cellular types: CD3, CD4, CD8 and FOXP3 bind to T cells, CD20 to B cells and CD 57 to NK cells[46].

In the complex relationship between the host immune microenvironment and cancer occurrence and progression, TILs seem to gain bidirectional regulation abilities. In one way, tumor neoantigens captured by the DC are presented on MHC molecules to the T cells, leading to the activation of effector T cells, with subsequent infiltration of the tumor and destruction of the cancerous cells. In addition, these activated cells secrete inhibitory cytokines, with the augmentation of antitumor effects[47,48]. On the other hand, TILs may help cancer to proliferate, either by creating an appropriate environment for tumor growth or by protecting tumor cells to survive[49].

The *stromal TILs* represent the mononuclear inflammatory cells infiltrating tumor stroma, whereas *intratumor TILs* define the intraepithelial lymphocytes/mononuclear cells within the tumor. Stromal TILs were shown to predict disease free survival (DFS) of patients[50].

The assessment of TILs as a prognostic biomarker in gastric cancer patients has led to controversial conclusions. The presence of various subsets of cells seems to differently influence the patient`s prognosis. Studies have shown that high density of intratumor TILs are associated with better prognosis (HR = 0.55)[51-53]. Additionally, some data in the literature revealed that an increased number of CD8+T cells, both intra- or extra tumor located, is associated with an improved DFS and OS[54-57]; in contrast, the results of a recent study showed that an increased number of CD8+ cells correlate with poor overall survival and increased expression of programmed-death ligand 1 (PD-L1)[58]. Other studies showed that a high density of intratumor FOXP3+ T reg is correlated with a poor OS, whereas an extratumor high density of this cell type leads to an increased OS[59-64]. The increased intratumor Treg/CD8+ lymphocytes ratio is correlated with a decreased OS[65] and the presence of T helper17 and T helper22 with tumor progression[66]. Moreover, an increased T helper1/T helper2 of CD4+ T lymphocytes represents a favorable prognostic factor[67]. A better OS was associated with an increased intratumor presence of various immunocytes, such as CD3+T cells[57,68], CD57 NK [51,57,69,70], CD45RO+ (memory T cells)[71], and T-bet+ (marker for T helper 1 lymphocytes)[72]; in addition, an increased DFS was observed in the case of high intratumor density of CD20 (surface marker of B cells)[73]. The data show a decrease in CD3+TILs density along with tumor progression[74]. On the other hand, the subgroup of CD45RO+ T lymphocytes seems to prevent peritoneal spreading of gastric neoplasias[75]. All the data from the literature demonstrate that high densities of CD8+, CD3+, and CD57+TILs and low densities of FOXP3+ TILs represent favorable prognostic factors in gastric neoplasia.

CD4+T cells secrete various cytokines, such as IL-17, the role for which the data from the literature reveal controversial results. Some of the studies showed that this cytokine could stimulate tumor angiogenesis, growth, and spreading[76], while other studies show that IL-17 exhibits anticancer effects, either by stimulating the cytotoxic activity of TIL[77] or by stimulating the maturation of DC [78].

The study of Yuan *et al*[79] revealed that gastric cancers present an increased percentage of CD4+T lymphocytes and lower CD8+T cells (with an increased CD4+/CD8+ ratio) compared to blood and, further, to paraneoplastic tissue. In addition, the number of TILs of effector and memory T cell type is significantly higher than in case of circulating T cells. As we have already mentioned, the involvement of Tregs in antitumor immunity is controversial, and their role may differ according to the type of cancer[80,81]. These authors considered CD4+CD25 highCD127low as being the most specific marker to define the Treg population, with the percentage of these cells being increased among TILs, demonstrating the accumulation of immunosuppressive Tregs at the site of a gastric tumor[79]. Recent studies show that the co expression of PD-1+ and Tim-3+ define the most hypo-functional T lymphocytes[32,82]. The percentage of these cells among TILs was significantly increased in gastric tumors, especially in patients with advanced stages, suggesting that they may be implicated in a tumor immune escape phenomenon and that TILs in gastric cancer show T cell dysfunction. The combined blockade of these molecules seemed to have a synergistic effect of IFNγ production and, therefore, may provide new promising immune modulating strategies[79].

Finally, the strategies of immunotherapy in gastric cancer are directed on TILs and are based on augmenting anticancer immunity by blocking the interaction of CD8+T lymphocytes-related receptors such as CTLA-4 and PD-1 and their ligands situated on tumor cells (PD-L2 and PD-L1). Furthermore, it may include the stimulation of the local immune response by targeting and modulating different subsets of CD8+ TILs.

***Significance of tumor infiltrating DCs in gastric cancer***

Immunotherapy needs the activation of the cellular immune responses following the presentation of the tumor antigen peptides by DCs to T cells[83].

It has been demonstrated that the density of tumor infiltrating DCs correlates with the staging and prognosis in gastric neoplasias, and patients with a high amount of DCs present an improved OS compared with patients with a lower density of DCs[69]. The study of Tsujitani et al showed that the use of postoperative adjuvant immunotherapy exhibits beneficial results on the survival of gastric cancer patients with reduced tumor DC infiltration[84].

***Classification of the immune microenvironment of the gastric cancer according to the presence of CD8+TILs and PD-L1 expression***

Based on the existence of tumor-infiltrating lymphocytes (TILs) and PD-L1 expression, it has been proposed to categorize the tumors into four types[85] as follows: type I (TILs+PD-L1+), presenting adaptive immune resistance; type II (TILs- PD-L1-), revealing immune ignorance; type III (TILs-PD-L1+), showing intrinsic induction; and type IV (TILs+ PD-L1-) in which other suppressors have a role in initiating immune tolerance. This stratification may have a certain prognostic role and may guide efforts towards a specific immunotherapeutic strategy.

Because tumor response to immunotherapy using PD-1 blockade requires the presence of CD8+TILs that are downregulated by PD-1/PD-L1 activity[86], this type of therapeutic strategy in type I cancer might improve prognosis. As tumors included in type II and III lack TILs, the combination of immune checkpoint inhibitors with a vaccine that is capable to induce T-cell activation, migration and infiltration at the tumor site might improve the clinical outcome in these patients[87].

***Correlation between TILs and gastric cancer subtypes***

Among the four molecular subtypes of gastric cancer, EBV(+) and MSI tumors often show the activation of immune mechanisms, being associated with a high density of TILs, which has been correlated with an improved cancer specific survival[88]. An increased number of CD8+ and FOXP3+ TILs were associated with improved OS in MSI-H gastric cancers[55] and EBV-associated cancers[87]. In addition, another study showed that a high number of CD3+ and CD8+ TILs in EBV-associated and MSI gastric cancer subtypes are associated with better survival[56].

Advanced TNM stages of EBV(+) tumors were correlated with a reduced density of CD4+, CD8+ and Foxp3+ TILs and PD-1 expression. Additionally, PD-L1 expression was shown to predict a reduced survival in EBV-associated cancers. Approximately two-thirds of EBV(+) gastric cancers were proved to present a type I or IV microenvironment associated with a better prognosis by inducing adaptive immune responses (type IV showed the best 5-year OS), whereas more than 70% of negative EBV tumors belong to the type II and III microenvironment, showing an absence of an immune response and a poor prognosis[87].

All these results are indicating the possibility of different subsets of TILs to be used as prognostic markers in these specific categories of patients. Moreover, EBV-associated and MSI gastric cancer categories might become potential targets of immunotherapy.

***Significance of peripheral immune status***

Myeloid-derived suppressor cells (MDSCs) represent immune suppressive cells with the ability to inhibit T cell (CD4+ and CD8+) activation and increase T cell apoptosis[89]. The data showed that high intratumor density of MDSCs is related to a poor prognosis in gastric cancer[80,90]. Peripheral blood granulocytes-MDSCs are significantly increased in cancerous patients[91]. Shoji *et al*[92], noted that an increased proportion of granulocytes-MDSCs prior to chemotherapy represents a negative prognostic factor for PFS in advanced gastric cancer patients receiving cisplatin-based chemotherapy and tends to be associated with poor OS.

Several papers showed that IL-8 is involved in gastrointestinal carcinogenesis by its ability to recruit MDSCs[93]. In addition, an increased number of circulating MDSCs was associated with advanced tumor stages, increased serum IL-8 levels, and dysfunction of T cells[94]. In this light, IL-8 seems to determine an adverse immune status[92].

Currently, numerous papers target defining the host inflammatory response to cancer. Due to a release of cytokines, the systemic inflammatory response seems to be responsible for the promotion of angiogenesis, DNA alteration, and tumor proliferation[95].

Studies have demonstrated that the neutrophil to lymphocyte ratio (NLR) has a prognostic value in patients with solid cancers[96-98]. In oncologic patients, lymphopenia is a marker of deficient cell‑mediated immunity, while neutrophilia is a sign of response to systemic inflammation. Furthermore, malnutrition/ hypoalbuminemia show correlation both with immune suppression and systemic inflammation, which are phenomena overexpressed in advanced tumor stages[99,100]. Gonda et al consider that NLR might be a useful biomarker for assessing tumor response to chemotherapy, immune suppression, malnutrition and unfavorable prognosis in gastric cancer patients. Moreover, it has been suggested that combining anti-inflammatory agents with chemotherapy provides an enhanced efficiency of the treatment[101].

**History of immunotherapy**

The concept of immune modulating treatment was used for the first time by Edward Jenner (1798), who demonstrated that inoculating humans with cowpox could prevent smallpox occurrence. Over time, this strategy was implemented in developing serum and vaccinations. The efficacy of immunotherapy in cancer treatment was demonstrated by WB Coley (1891) who, by injecting streptococcal germs into a patient with unresectable cancer, determined the decrease of the tumor size[102]. Furthermore, Ehrlich (1909) has suggested that the host immune system could suppress the tumor growth by recognizing cancerous cells as non self. Half a century later, Burnett has proposed the theory of tumor immune surveillance[35], recently completed by Schreiber et al. with the concept of cancer immunoediting[49].

In recent years, many immunotherapeutic strategies have been developed, including treatments using monoclonal antibodies, cytokines, cytotoxic cells, T cells infusions and gene transferred vaccines[36], having the aim of either increasing the host antitumoral response capacity or increasing the immunogenicity and susceptibility to treatment of the tumor cells[103].

**Current immunotherapeutic strategies in gastric cancer**

The development of an effective immunotherapeutic management for digestive cancers has evolved relatively slow, with the majority of these immune modulating approaches still under assessment in early phase clinical trials, mostly because of their well-known lack of antitumor effector T lymphocytes responses and their decreased immunogenicity[104]. Despite these specific aspects, some immunotherapeutic approaches, such as those using anti PD/PD-L1 immune checkpoints inhibitors, proved to be effective also in cancers defined by a poor immunogenic nature[105].

Immunotherapeutic strategies may be classified into[42]:

**Active immunization strategies,** **including:** (1) adoption of cytokines (e.g., IFNγ, IL-10, IL-2) - to date, leading to inconclusive results[103]. (2) vaccination strategies that include: vaccines using peptides/proteins recognized by CD8+ and CD4+ lymphocytes, such as various tumor-rejection antigens (including melanoma-associated antigen (MAGE-3) or HER-2/neu)[106]; new vaccines using a compound formed by an immunogenic protein fused with peptide (Z12) that determines a persistent antitumor T-cell response[107]; DC-based vaccines; RNA-based vaccines, *etc*. (3) immune checkpoint inhibitors (anti-CTLA4, anti-PD/PD-L1). (4) combination of different immunotherapeutic strategies. And (5) combination of immunotherapy with standard treatment

**Passive immunization strategies:** adoptive cell therapy (ACT) using TILs - refers to the passive transfer of antitumor T lymphocytes into a tumor-bearing host followed subsequently by the direct destruction of this cancer[108].

Current immunotherapeutic strategies are presented in Figure 2.

***Cellular immunotherapies in gastric cancer***

Currently, immunotherapy in gastric cancer patients includes cell-based strategies aimed either to activate cytotoxic T lymphocytes directed against cancer cells or to bind molecules expressed by tumor cells.

**ACT:** This technique refers to injection of different tumor-specific T lymphocytes into a cancer patient, such as cytokine- and anti-CD3 monoclonal antibody- induced killer cells, as well as TILs.

(1) Cytotoxic T-cell therapy

In a preclinical study, cytotoxic activity of peripheral blood lymphocytes extracted from gastric tumor patients or from healthy individuals was induced using different HLA-A matched allogeneic gastric cancer cells, exhibiting antitumor efficacy against HLA-A2 and HLA-A24 gastric cancer cell lines[109]. Furthermore, Kawamoto *et al*[110] demonstrated the efficiency of peptide-based immunotherapeutic strategies by proving that cytotoxic T lymphocytes were able to kill HLA-A-0201/2402 colon and gastric cancer cells, which were positive for mitotic centromere-associated kinesin (MCAK) (a new cancer antigen) in an HLA-I restricted way. In addition, MHC-1 restricted T cells were obtained from primary tumors, metastatic lymph nodes, and ascites of autologous gastric cancer and were proved to detain different recognition characteristics towards gastric cancer antigens[111]. Moreover, splenic MAGE-specific cytotoxic T lymphocytes against HLA-A2 cancer cells antigen, existing in testis and several neoplasias (including gastric cancer), were successfully obtained and tested[112].

Another preclinical study has revealed that cytokine- induced killer (CIK) cells, mostly by stimulating IFN- γ and tumor necrosis factor-alpha, exhibit antiproliferative effects against the MGC-803 gastric cancer cell line[113]. In addition, Kim et al demonstrated the benefits of using ACT with CIK cells in gastric cancer patients. These CIK cells, isolated from the human peripheral blood mononuclear cells and activated by IL-2 and anti-CD3 antibody, were able to suppress the MKN74 human gastric cancer cell line in vitro and inhibit tumor proliferation in a nude mouse model[114].

As gastric cancers usually show paucity of stroma infiltration, *in vivo* studies have suggested administration of immunotherapy using ACT combined with chemotherapy[115]. Furthermore, the chemotherapeutic drug oxaliplatin, by stimulating high-mobility group box 1 protein to induce anticancerous T lymphocytes, is capable of producing an immunogenic cancer cell death[116]. In several *in vitro* and *in vivo* studies on drug-resistant gastric cancer, a high amount of cytokines was induced by combining this drug with CIK cells[117]. It was suggested that chemotherapy associated with T lymphocytes reduction would be capable of enhancing the results of ACT therapy by stimulating the persistence of endogenous T cells in circulation, while depleting autoimmune reactions on healthy tissues but unfortunately, these results occurred at the expense of severe infectious adverse events in these patients[118].

Moreover, the results obtained by combining ACT treatment with an antibody directed against both anti-epidermal growth factor receptor (EGFR) and anti-epithelial cell adhesion molecule (EpCAM), that specifically targets the simian virus 40 (SV40) T antigen-specific T cells (previously transduced with a truncated human EGFR), showed a better tumor reduction and OS than with ACT alone [119].

Du *et al*[120] used a mouse model of gastric cancer and concluded that ACT by peritoneal injection of CIK might be both a beneficial and minimally invasive strategy for treating this type of cancer.

The first clinical trial proving the beneficial results of ACT in humans used lymphokine-activated killer cells plus IL-2 in patients with metastatic melanoma, leading to the approval of the treatment for this group of patients[121-123]. Furthermore, this strategy determined a significant tumor reduction in patients having different tumors[124].

Zhang *et al*[125] showed that administration of expanded activated autologous lymphocytes, which were stimulated by anti-CD3 mAb and IL-2, to gastric neoplasia patients led to a prolonged OS compared to the group that received only standard chemotherapy.

The efficiency assessment of combined treatment using CIK cells and chemotherapy as adjuvant therapy in stage II/III of gastric neoplasia after curative gastric resection revealed a significant prolonged OS *vs* the group using chemotherapy alone[126]. In a similar context, a clinical trial analyzed the possible toxicities of adjuvant ACT associated with chemotherapy (including 5-fluorouracil/capecitabine), showing an improvement in both DFS and OS, without the development of severe adverse reactions[127]. Moreover, a clinical trial assessing this type of combined adjuvant treatment in stage III/IV (M0) gastric cancer patients after R0/D2 resection, showed a significantly longer 5-year OS and DFS *vs* adjuvant chemotherapy[128].

In cases of advanced gastric cancer, several clinical studies have also proved an increased response rate, better quality of life and even an increased OS in patients treated with chemotherapy (FOLFOX4) plus CIK cells *vs* chemotherapy alone[129,130].

By assessing the administration of a chemotherapeutic regimen followed by autologous CIK cells, the results highlighted an improved remission rate in gastric cancer patients, associated with tolerable and reversible side effects. Standard chemotherapy using XELOX plus CIK cells administered intraperitoneally in gastric cancer patients showed a marked decrease in ascites volume and OS prolongation[131].

In a pilot study, patients with advanced gastric cancer who received gamma delta T cells with zoledronate intraperitoneally as a local treatment for carcinomatous ascites, showed a significant reduction both in the number of peritoneal tumor cells and ascetic volume with no serious side effects[132].

Several clinical trials are currently investigating the tumor responses after adjuvant administration of ACT plus chemotherapy after surgical resection in advanced gastric cancer patients[133]. In a phase II clinical study involving gastric cancer patients in stages I-III, the adjuvant combination of autologous tumor lysate-pulsed dendritic and CIK cells (Ag-D-CIK) plus chemotherapy is currently being evaluated following curative resection[134].

A clinical trial is currently investigating the safety and efficiency of infusing chimeric antigen receptor (CAR) T cells specific for EpCAM into relapsed or refractory gastric cancer patients[135].

A phase I/II clinical trial is currently investigating the benefits of infusing CAR T cells targeting mucin 1 (MUC1) in several solid cancers (including gastric tumors), as its overexpression leads to chemotherapeutic-refractory tumors[136].

A phase Ib clinical trial on advanced gastric cancer expressing CEA, assesses the efficacy of injecting anti-CEA CAR T cells into the hepatic artery targeting liver metastasis[137].

Additionally, a phase I/II clinical study on HER2(+) gastric cancer patients (defined as HER2 in immunohistochemical tumor tissue greater than or equal to 2 levels) with liver metastasis is analyzing the cytotoxic potency of engineered pluripotent stem cells and T cells, which specifically bind to HER2[138]. In addition, another phase I clinical trial assesses the safety profile of administrating autologous T cells equipped with a bi-specific antibody (HER2 Bi-Armed T cells) in gastric and esophageal neoplasias[139].

Patients with metastatic gastric cancer are also investigated regarding a combination of S-1 plus dendritic cell activated CIK (DC-CIK) (phase I/II clinical trial)[140].

(2) Adoptive immunotherapy using TILs

The use of ACT with TILs is not associated with an immediate effect because this therapeutic protocol requires approximately six weeks, as T cells must undergo the following preparation steps before infusion: first, they are isolated from tumor tissue; next, they are *in vitro* expanded; and finally, tumor-specific T cells are selected [141].

The immunotherapy using TILs has led to encouraging results in preclinical models[142] but not in all clinical studies (except for melanoma)[143,144].

In most gastric cancer patients, TILs exhibit a specific type-1 T-cell response to cancer antigens. It is important to note that in order to obtain *“in vivo”* destruction of the tumor cells, the efficacy of tumor-specific T cells usually needs to be enhanced by combining vaccination using specific cancer antigens/peptides or by injecting *in vitro* expanded autologous cancer-specific T cells [103].

Moreover, TILs can sometimes stimulate proliferation of tumor cells. Studies show that HP0175-specific TILs in gastric cancer patients infected with Helicobacter-pylori (Hp) determine gastric Th17 response, exhibiting a pro-inflammatory low cytotoxic TIL response; Th17 cells promote tumor progression through the promotion of inflammation by secretion of IL-17 and other interleukins, which could induce proliferation and migration of cancer cells; therefore, TILs reveal a correlation between Hp infection and gastric cancer development[145].

Because studies have demonstrated that in the tumor areas, a high Tregs/CD8+ ratio represents an independent factor for poor OS, a combination of the deletion of Tregs plus the stimulation of effector T cells may represent an effective immunotherapy in gastric cancer patients[65].

ACT using TILs isolated from the patient’s tumor was also assessed in cases with gastric neoplasia[146]; the results of a clinical trial showed a longer OS using a combined treatment of ACT using TILs and standard chemotherapy *vs.* chemotherapy alone[143].

(3) Adoptive immunotherapy using NK cells

The data show that NK cells exhibit cytotoxic activity against allogeneic and autologous cancer cell lines, including gastric cancer cells lines[147] and could prevent tumor metastatic spreading[148,149]; additionally, intra-tumor infiltration of NK cells is associated with a longer survival in neoplastic patients[150]. Patients with advanced gastric adenocarcinoma having a high density of NK cells demonstrated a prolonged survival rate *vs* those with the low density NK[151]. The number of apoptotic NK cells (Fas+ NK cells) is significantly higher in gastric neoplasia patients *vs* normal controls and is correlated to the tumor progression[152].

Clinical data revealed a favorable prognostic role of NK cells in gastric cancer patients, with a high level of CD57 antibody expression in gastric tumors correlated with a reduced size of tumors, N0 tumors, more surgical resections and prolonged 5-year OS[51].

By culturing autologous peripheral blood mononuclear cells with K562 cells, researchers were able to obtain cytotoxic NK cells from cancer patients[147], suggesting a possible role of immunotherapy using autologous expanded NK cells in clinical practice. In this regard, a clinical study using a combination of cell-based immunotherapy with autologous NK cells, γδ T cells, and CIK cells plus chemotherapy showed a statistically significant improvement of the two-year progression free survival and quality of live, but without demonstrating a significantly prolonged OS in gastric cancer patients[153].

The safety and efficacy of therapy with trastuzumab and NK cells in the treatment of gastric cancer is currently assessed in a clinical trial[154].

It has been demonstrated that lupeol, which exhibits a curative effect on various diseases, has also the ability not only to stimulate the proliferation and the cytolytic activity of NK cells against gastric tumor cells but also to inhibit the proliferation of some gastric cancer cell lines. Therefore, this agent might be included, either alone or in combination with ACT using NK cells, in the therapeutic strategies of gastric tumors[155].

***Cancer vaccines***

Cancer vaccines have the aim of activating and increasing the number of effector T lymphocytes, leading to the augmentation of existing immunity, development of novel immunological response, and therefore an improved anticancer immunity[36].

Peptides derived from tumor-associated antigens (HER2/neu-derived peptide[156] and MAGE[157] are captured by antigen presenting-cells (such as DC)) and presented by means of MHC type I for presentation to cytotoxic T cells, and by means of MHC type II for presentation to T helper cells, determine their activation, followed by the destruction of tumor cells.

There are several types of vaccinations that have shown promising results in gastric cancer patients:

(1) HLA-A∗2402-restricted URLC10-A24-177 and vascular epidermal growth factor receptor (VEGFR1-A12-9 1084) epitope peptide cancer vaccines in advanced chemotherapy-resistant gastric cancer patients – these are safe and induce enhanced specific cytotoxic T cell immune response[158].

(2) vaccination using survivin epitope peptide – a recent study suggested an excellent efficiency in gastric cancer patients upon inducing survivin-derived peptide-specific cytotoxic T-lymphocytes from mononuclear cells isolated from blood of healthy individuals[159].

(3) autologous gp96 vaccine in addition to chemotherapy as an adjuvant treatment in patients with resected gastric cancer (phase II study) - glycoprotein (gp)96, belonging to the group of autologous tumor-derived heat shock proteins (HSPs), binds tumor-associated antigens, constituting the HSPs–peptide complexes that promote the activation of APCs, as well as the release of various cytokines, enhancing T-cell antitumor immune response; the vaccination seemed be well-tolerated and is associated with a potential for prevention of gastric cancer recurrence[160].

(4) trials using DCs-based anticancer vaccines - *ex vivo* expanded DCs were able to generate antigen-specific T lymphocytes responses both in animal models[161], and in clinical trials[162]: (1) vaccination using dendritic cells pulsed with HER-2/neu peptide generated tumor shrinkage (phase I study) [143]; (2) vaccination using DC pulsed with nanoparticles MAGE-3 peptide–loaded stopped tumors from proliferating in a mouse model of gastric cancer (phase I study)[163]; and (3) vaccination with cancer-loaded autologous DCs (stimulated by autologous apoptotic gastric tumor cells) determined the activation of memory T cells[164].

**RNA-based DC vaccines:** By using stabilized mRNA, DCs transfected with mRNA coding for tumor-associated antigen/whole tumor RNA were able to generate potent immune responses both in mouse models[165,166] and clinical studies involving melanomas and renal cell carcinomas[167,168].

These RNA-based vaccines seem to present some potential benefits over the classical vaccination techniques: they are pharmaceutically safer because of the presence of transient and cytosolic active mRNA (lack of genomic integration); they have the ability to target multiple tumor-associated antigens; they are not associated with severe adverse events; and they are not MHC-restricted[36].

Because the clinical efficiency of DC vaccines is limited because of the short survival of DCs, mostly due to the cytolytic properties of DC-activated CD8+ T cells[169], their efficiency has been increased by using small interfering RNA (siRNA)-targeting phosphatase and tensin homolog (PTEN), which is involved in negative feedback in the signal transduction of the PI3K/AKT pathway[170]. This technique has increased the number of cancer-specific cytotoxic T cells and the antitumor immune response[171].

The results were improved by targeting the immunosuppressive IL-10 receptor in association with the siRNA DC vaccine[172], by using GM-CSF gene-modified DC[173], and by removing Tregs along with DC vaccination[174].

**Strategies of combining vaccination with chemotherapy:** The combination of an adjuvant Bacille Calmette-Guérin (BCG) vaccine with chemotherapy *vs.* chemotherapy alone resulted in a better OS in patients with radically resected locally advanced gastric cancer[175].

Vaccination using gastrin-17 diphtheria toxoid (G17DT)-targeting gastrin peptide combined with chemotherapy (cisplatin plus fluorouracil) increased the OS of patients with advanced gastric cancer (phase II clinical trial)[176].

Vaccination using peptides derived from human VEGF receptors 1 and 2 combined with standard chemotherapy (S1+cisplatin) improved the OS in patients with advanced gastric cancer[177].

Literature data show that vaccination is a safe and tolerable strategy that is associated with better prognosis in gastric cancer patients, especially when it is performed in addition to classical chemotherapy.

***Immunotherapy using checkpoint inhibitors***

Immune checkpoints represent inhibitory pathways that are critical for maintaining self-tolerance and physiological homeostasis by controlling the intensity of physiological immune responses to prevent tissue injury, particularly when the immune system is fighting an infection. Additionally, they may also allow immune escape of cancer cells[36].

Immune checkpoint molecules, such as cytotoxic T lymphocyte protein-4 (CTLA-4) and programmed cell death protein-1 (PD-1), are involved in the inhibition of T-cell activation via different pathways.

CTLA-4 is a co-inhibitory molecule exhibited on activated T lymphocytes and T regulatory cells, whose receptor on T cells interacts with its B7-1/B7-2 ligands located on antigen-presenting cells (APCs), subsequently suppressing the T-cell stimulatory signal mediated by CD28[178]. CTLA-4 expression is stimulated only in the context of T-cell activation; afterwards, it competes with CD28 to bind to B7 molecules and decrease the immune response. By inhibiting this interaction using an anti-CTLA-4 antibody, T-cell activation and proliferation is promoted, along with a decrease in immunosuppressive Treg cells among TILs[179].

PD-1 represents a co-inhibitory receptor that is found on the surface of several types of cells, such as activated T cells, Treg cells, and monocytes. It has two ligands, PD-L1 and PD-L2. PD-L1 is expressed on both immune and tumor cells, while PD-L2 is mostly expressed on APCs. In tumors, PD-L1 that is expressed on tumor cells binds to PD-1 on activated T cells that reach the tumor and generates a suppression signal for the activation of T cells, which become unable to destroy tumor cells, leading to a decrease in both cellular and humoral immune responses[180,181]. Unlike CTLA-4, which is considered to be necessary for T-cell activation, the PD-1/PD-L1/2 pathway seems to protect tumor cells from attack by T lymphocytes. It has been demonstrated that by inducing antibody-mediated blockage of the PD-1/PD-L pathway followed by the inhibition of this checkpoint, treatment is able to enhance the anticancer immune response of the host[22,182].

Several genetic studies have shown a possible correlation between PD-1 or CTLA-4 polymorphisms and the development of gastric cancer[183-185]. Additionally, the role of CTLA-4 gene promoter hypermethylation has been demonstrated as a risk factor for gastric tumor development, with CTLA-4 expression being significantly higher in gastric tumor samples *vs.* normal tissue[186].

**Prognostic significance of PD-1/PD-L1 and CTLA-4 expression in gastric cancer:** Saito *et al*[187] showed that PD-1 expression in CD8+ and CD4+ T cells in gastric tumors was significantly higher *vs.* in normal gastric mucosa. Wu *et al*[188] showed that PD-L1 expression was encountered in 42% of gastric cancer tissues but not in normal gastric mucosa.

A meta-analysis by Gu et al that was based on 15 studies (most of them conducted in Asia) including 3291 patients, showed a high variability of PD-L1 immunohistochemical expression among studies, ranging between 14.3% and 69.4%, due to the differences in cut-off values (between > 1% and > 50%)[189].

Unlike in other tumors, such as lung cancer or melanoma, there is scattered PD-L1 expression in gastric cancer cells, which mostly occurs in infiltrating myeloid cells at the tumor invasive front[25,58].

There are several papers suggesting that PD-L1 expression in tumor cells may be upregulated by genomic alterations and oncogenic signaling, either through the phosphatidylinositol-3-kinase-protein kinase B (PI3K-AKT) or signal transducers and activators of the transcription (STAT) 3 pathway. Moreover, PD-L1 is upregulated by the microRNA-200/zinc-finger E-box-binding home-obox 1 (ZEB-1) axis, which is closely related to epithelial-mesenchymal transition (EMT) conversion and to IFN-γ produced by TILs[190-192]. Mimura et al. showed that membranous PD-L1 immunohistochemical expression in gastric tumor cells was significantly correlated with the number of CD8+ TILs and IFN-γ positive cells in the tumor. Additionally, an enhanced cytotoxic effect of anti-PD-L1 monoclonal antibody treatment was observed after prior IFN- γ exposure[193].

Literature data regarding the expression of PD-1/PD-L1 as a prognostic in gastric cancer showed controversial results. Dai et al. reported an association between PD-L1 expression and an increased density of TILs; this increased density of TILs and higher levels of PD-L1 mRNA in gastric cancer was signiﬁcantly correlated with a better prognosis[53]. Some studies showed a significantly improved prognosis in patients with PD-L1 positive tumors[194,195]; conversely, others reported an association between PD-1/PD-L1 expression in cancer cells and TILs and advanced tumors, increased tumor size, the presence of deep invasion, lymph node metastasis, and perineural invasion and a significantly worse prognosis in these patients[196-198]; and the third group of papers found no influence of PD-L1 on the prognosis of gastric cancer[199].

A few recent meta-analyses have shown a correlation between PD-L1 and gastric cancer prognosis, demonstrating that PD-L1 overexpression is a worse prognostic factor in these patients[200-202]. Additionally, Zhang *et al*[203] performed a recent meta-analysis that included 10 studies with 1901 gastric cancer patients and showed that PD-L1 expression was associated with a shorter OS and a poor clinicopathological status.

Fang *et al*[204] highlighted that PD-L1 was expressed both in tumor cells and in TILs. PD-L1 positivity in tumor cells was associated with differentiation, while its expression in TILs was correlated with a late stage of the disease, no surgery and the OS. Patients with PD-L1 + TILs had a significantly poorer five-year OS than those without PD-L1 expression (14.2 *vs* 18.3; *p* = 0.001). A study by Gao et al showed that PD-L1 and PD-L2 positivity in primary tumors and metastatic lymph nodes, decreased the number of CD8(+) T cells and the amount of PD-1 positive expression on CD8(+) T cells in primary tumors were prognostic factors that were correlated with a poor prognosis in stage II/III gastric cancer patients[205].

Again, the results of the meta-analysis by Gu and collaborators[189], showed that gastric cancer patients with deeper tumor infiltration, lymph node metastasis, venous invasion, and EBV+ and MSI subtypes were more likely to be PD-L1 positive. Moreover, for the subgroups of Asian patients and patients with stage II/III gastric cancer, cut-off values greater than 50% and cytoplasm/nuclear PD-L1 expression within tumor cells were positively associated with the OS. The results of this meta-analysis demonstrated that PD-L1 overexpression represented a significantly adverse prognostic factor in gastric cancer, fitting the theory of the cancer-immunity cycle[47].

Many other studies[32,206] have also demonstrated that EBV+ and MSI gastric tumors tend to be positive for PD-L1 expression. These subtypes have a rich infiltration of lymphocytes, especially CD8+T cells, in the tumor stroma; therefore, they may be categorized as medullary carcinomas. Moreover, in this context, PD-L1+ expression is associated with a significant increase in the number of CD8+T cells at the tumor invasive front and with the ability of immune cells to infiltrate the center of the tumor[32]. Both EBV+ gastric cancers and MSI+ gastric cancers have IFN-γ response genes, therefore, PD-1 pathway signaling seems to be a crucial mechanism for controlling a previous cytotoxic anticancer immune response[189]. The EBV+ subtype is associated with amplification of the 9p24.1 locus, which harbors the PD-L1/PD-L2 genes; PD-L1 positivity is found in 50% of the cancer cells and 94% of immune cells in this subgroup[194,207]. Additionally, approximately 33% and 45% PD-L1+ expression levels are encountered on tumors cells and immune cells, respectively, in MSI gastric tumors[208]. These data suggest that EBV+ and MSI gastric cancers may be preferred candidates for PD-1 blockade immunotherapy. These types of neoplasias that are associated with rich inflammatory infiltrates are considered “hot” or inflamed tumors, while poorly immunogenic cancers are termed “cold.” “Hot” tumors are characterized by the expression of immune-inhibitory signals, such as PD-L1, indoleamine-2,3-dioxygenase (IDO), and Treg cells[209,210], which counterbalance the effects of cytotoxic T lymphocytes. Therefore, by using combined treatments to convert cold into hot tumors may increase the proportion of patients who benefit from immunotherapy[211].

While some data from non-small cell lung cancer (NSCLC) and other cancers have revealed that PD-L1 immunohistochemical positivity in cancer and/or in immune cells (bioptic specimens) is correlated with beneficial results after checkpoint inhibitor immunotherapy using monoclonal antibodies[212], other studies have shown tumor responses to PD-L1 therapies in tumors with PD-L1 (-) cancer cells[213].

The location of PD-L1 expression in TILs situated at the invasive front or even at the tumor center may negatively influence its use as a biomarker. Furthermore, stromal rather than membranous expression of PD-L1 may be the cause for the slightly poorer responses to single-agent PD-1 checkpoint inhibitors in gastric cancer *vs* other tumor types[208].

Because there are variations in the techniques, antibody clones used and cutoff values for PD-L1 positivity that limit cross-trial comparisons of different tumor types, including gastric cancer, some researchers have proposed harmonization of PD-L1 testing to standardize the results. Recently, Jiang *et al*[214] elaborated an immunohistochemical-based immunoscore on a cohort of 879 Chinese gastric cancer patients and demonstrated that a high immunoscore corresponded to lower recurrence rates and improved OS after adjuvant therapy.

Data suggest that there is racial and geographical variability in the tumor-immune microenvironment that is related to different responses to immunotherapy; for example, non-Asian gastric cancer patients have tumors that are rich in TILs and are associated with high CTLA-4 signaling[215].

The study by Schlößer et al. found positive CTLA-4 expression in the tumor microenvironment of 86% of gastric cancer patients that was correlated with poor OS[198].

**Clinical trials using checkpoint inhibitors:** Clinical trials using CTLA-4 and PD/PD-L1 checkpoint inhibitors are listed in Table 1.

(1) CTLA-4 checkpoint inhibitors

Tremelimumab

Tremelimumab (formerly ticilimumab, CP-675,206) is a fully human [monoclonal antibody](https://en.wikipedia.org/wiki/Monoclonal_antibody) against CTLA-4 that received FDA approval for the treatment of mesothelioma. It was the first immune checkpoint inhibitor that was investigated in patients with gastroesophageal tumors.

A phase II study[216] investigated tremelimumab (at a dose of 15 mg/kg every 90 d, which is presently considered sub therapeutic) in 18 patients with advanced esophageal, gastroesophageal junction or gastric adenocarcinoma. Although the median time to progression and OS were relatively short, at 2.83 and 4.83 mo, respectively, approximately one-third of patients were alive at one year. Currently, there is an ongoing clinical trial assessing whether tremelimumab is associated with the anti-PD-L1 antibody durvalumab; the dose of tremelimumab used as a monotherapy is 10 mg/kg every 4 wk[9].

Ipilimumab

Ipilimumab (trade name Yervoy, previously known as MDX-010 and MDX-101) was approved by FDA for the treatment of patients with unresectable/metastatic melanoma, after at least one line of systemic treatment has been performed (2011)[217]; the indications for this drug were recently extended to include pediatric patients (12 years and older) (2017) and adjuvant treatment of patients with cutaneous melanoma with pathologic involvement of regional lymph nodes greater than 1 mm in size who have undergone complete resection (including total lymphadenectomy); in addition, it was recently approved as the first-line treatment, in combination with nivolumab, for patients with intermediate or poor risk with advanced renal cell carcinoma (2018).

A randomized phase II trial enrolled patients with unresectable, locally advanced/metastatic gastric or gastro-esophageal junction cancer with partial response/stable disease after first-line chemotherapy with a combined fluoropyrimidine plus platinum regimen to receive either the best supportive care (consisting of continuation of fluoropyrimidine) or ipilimumab. The primary end-point of the study was immune-related PFS. Unfortunately, the study was ended earlier due to the lack of clinical efficiency of ipilimumab. The PFS was shorter, and the toxicities were higher in the ipilimumab group, with similar OS for both groups[218].

(2) PD-1 and PD-L1 checkpoint inhibitors

Anti-PD-1 antibodies - Nivolumab

Nivolumab is a humanized immunoglobulin G4 monoclonal antibody (mAb) that targets PD-1 and has shown efficacy in various cancer types. The FDA approved nivolumab for the treatment of recurrent/metastatic melanoma, metastatic non-small cell lung carcinoma/recurrent or metastatic squamous cell carcinoma of the head and neck (after progression on platinum-based chemotherapy), advanced renal cell carcinoma after antiangiogenic treatment, and Hodgkin lymphoma that has relapsed/progressed after autologous hematopoietic stem cell transplantation.

The phase I/II CheckMate-032 clinical trial analyzed the safety and efficacy of nivolumab as monotherapy or combined with ipilimumab in heavily pretreated patients with advanced/metastatic solid cancers, including gastric tumors (NCT01928394)[219]. The partial results assessing 160 patients included in 3 different arms (nivolumab 3 mg/kg (N3), nivolumab 1 mg/kg plus ipilimumab 3 mg/kg (N1+I3), and nivolumab 3 mg/kg + ipilimumab 1 mg/kg (N3+I1)) showed more frequent toxicities in the N1+I3 group. The most commonly encountered adverse events in all the subsets were fatigue, pruritus, nausea, diarrhea and loss of appetite; thyroid damage was more frequent in the N1+I3 group. The overall objective response rate (ORR) was 16%, which was the highest in the N1+I3 group 26%; the disease control rate (DCR) was of 38%. The median OS was longest in the N1+I3 arm (6.9 mo), followed by the N3 (5.0 mo) and the N3+I1 arms (4.8 mo). The response rates for patients with PD-L1+ tumors (>1% of immunohistochemically positive cells), both in patients treated with nivolumab monotherapy and combined treatment N1+I3 and N3+I1, were superior compared to patients with negative tumors (19% *vs*. 12%, 40% *vs.* 22% and 23% vs. 0%, respectively).

Because of the promising results obtained using N1+I3 combination, the CheckMate-649 phase III clinical trial (NCT02872116) was initiated. In this study, the clinicians plan to investigate 1349 patients with naive advanced or metastatic gastric/gastroesophageal junction (GEJ) cancer with both positive/negative PD-L1 expression to receive first line nivolumab plus ipilimumab or nivolumab plus standard chemotherapy (XELOX or FOLFOX) *vs* standard chemotherapy (XELOX or FOLFOX), having as primary endpoint the OS in patients with PD-L1+ tumors[220]. The data from the double-blinded, randomized, multicentric phase III trial ONO-12 (ATTRACTION 2) were presented at the ASCO Gastrointestinal Cancers Symposium 2017, demonstrating the benefits of nivolumab as a salvage treatment (third or later line) in patients with advanced unresectable or recurrent gastric or gastroesophageal junction cancer, either PD-1/L1 (+) or (-) *vs* placebo (NCT022673430). It was the first time that a prolonged OS was obtained for patients with heavily pretreated tumors using PD-1 inhibition[221]. There were 493 patients randomly assigned (2:1) to receive nivolumab (*n* = 330) or placebo (*n* = 163). The median OS was significantly increased with nivolumab *vs* placebo (5.26 mo *vs* 4.14 mo with placebo). The risk of death was lower in the nivolumab group *vs* placebo group (HR = 0.63; *p* < 0.0001); 68.5% of the patients in the nivolumab group died *vs.* 86.5% in the placebo group. Additionally, nivolumab treatment was associated with a significantly better OS rates at 6 and 12 mo compared to placebo (46.1% *vs* 34.7% and 26.2% *vs* 10.9%), Nivolumab treatment led to a longer median PFS (1.61 mo *vs* 1.45 mo) and higher ORR than placebo (11.2% with nivolumab *vs* 0%). Among the approximately 40% (*n* = 192) of patients with tumor samples, 12.3% in the nivolumab group and 16.1% in the placebo group had PD-L1 positive tumors. The analysis of PD-L1 expression status showed that median OS in patients with PD-L1 positive tumors was 5.22 mo in the nivolumab group and 3.83 mo in the placebo group (HR = 0.51). In patients with PD-L1 negative tumors, median OS was 6.05 mo in the nivolumab group, and 4.19 in the placebo group (HR = 0.72). Nivolumab resulted in an absolute survival benefit of 1.1 mo in median OS *vs* placebo. Nivolumab led to durable OS benefit that was sustained beyond 1 year *vs* placebo in heavily pretreated gastric cancer patients, regardless of PD-L1 expression status[222]. Because of the survival benefit demonstrated by the ATTRACTION-2 trial in this subset of difficult-to-treat gastric cancer patients, the approval of nivolumab in Japan was granted as a new treatment option that is beneficial for heavily pretreated advanced gastric cancer.

Anti-PD-1 antibodies - Pembrolizumab

Pembrolizumab (formerly MK-3475, trade name Keytruda) is a [humanized IgG4 isotype antibody](https://en.wikipedia.org/wiki/Humanized_antibody) that targets PD-1 receptor of lymphocytes. It has FDA approval for the treatment of unresectable or metastatic [melanoma](https://en.wikipedia.org/wiki/Melanoma), metastatic [non-small cell lung cancer](https://en.wikipedia.org/wiki/Non-small_cell_lung_cancer) (NSCLC) (certain situations), metastatic non-squamous NSCLC (PDL1+/-), as a second-line treatment for [head and neck squamous cell carcinoma](https://en.wikipedia.org/wiki/Head_and_neck_squamous_cell_carcinoma) (HNSCC), after [platinum-based chemotherapy](https://en.wikipedia.org/wiki/Platinum-based_antineoplastic), and for the treatment of adult/pediatric patients with refractory classic [Hodgkin's lymphoma](https://en.wikipedia.org/wiki/Hodgkin%27s_lymphoma).

The large multi-cohort, multicenter, nonrandomized, open-label phase Ib KEYNOTE-012 first demonstrated the efficacy of pembrolizumab as monotherapy (administered in a dose of 10 mg/kg every 2 wk or a 200 mg fixed dose every 3 wk) in PD-L1-positive recurrent or metastatic gastric or gastroesophageal cancer (NCT01848834)[25]. Of all the patients assessed, 39 patients (40%) had PD-L1-positive tumors, most of them being heavily pretreated patients. Treatment with single-agent pembrolizumab determined partial response (PR) in 22% and sustained disease (SD) in 13% of patients. The median PFS was 1.9 mo, with a 6-mo PFS of 26% and a median OS of 11.4 mo. The most common adverse events included loss of appetite, fatigue, pruritus, arthralgia and hypothyroidism. Pembrolizumab was well tolerated, with 13% of patients developing grade 3–4 toxicity. It is important to stress that approximately 60% of patients enrolled in this trial have previously received more than 3 lines of chemotherapy, representing a group of patients without any therapeutic response demonstrated by other studies. Almost two-thirds of the tumors revealed genomic profiling of a microsatellite-instability high (MSI- H) status. This genomic status correlates with high tumor mutational burden, and patients with MSI- H cancers, colorectal and non-colorectal, have developed encouraging responses to anti–PD1 therapy in various solid tumors[23]. This aspect remains to be established if present in gastroesophageal cancers. Additionally, in this study, tumor response correlated with an increased interferon gamma gene expression[223].

With the goal of improving immune treatment efficacy and having the knowledge that chemotherapy is capable of promoting immunogenic cell death[224], PD-1 inhibitors were associated with standard chemotherapy, which is a combined regimen proved to increase response rates in lung cancer[225]. The KEYNOTE-059 phase II clinical trial included 259 patients with advanced gastric or gastroesophageal cancer. Cohort 1 assessed the efficacy and safety of pembrolizumab monotherapy in patients with previously treated advanced gastric cancer. Approximately half the patients enrolled in cohort 1 had received 2 or more prior treatments for metastatic cancer, whereas others received 3 or more prior therapies. Among 259 patients, 148 (57.1%) were PD-L1 positive (by immunohistochemistry). Pembrolizumab elicited sustained ORR in 30 of 259 patients (11.6%) and complete response in 2.3%. These responses were observed irrespective of PD-L1 expression. The ORR was higher in patients with PD-L1 positive *vs.* PD-L1 negative tumors, 23 of 148 (15.5%) *vs* 7 of 109 (6.4%), respectively. A T-cell–inflamed gene expression profiling score was developed in this study, demonstrated to be significantly associated with pembrolizumab response; also, a significant nonlinear association was found between this score and PD-L1 expression. Of 174 patients (67.2%) assessed for MSI, 7 patients (4.0%) had samples that were MSI-high. It was observed a higher ORR in patients with MSI-high tumors than in patients with non–MSI-high tumors (57.1% *vs* 9.0%). However, prevalence of MSI high tumors was very low in this population (4%), and most responses were observed in non–MSI-high patients. Thus, in cohort 1, pembrolizumab monotherapy showed good responses and manageable toxicities after ≥ 2 prior lines of treatment[226,227]; these results led to an accelerated FDA approval of pembrolizumab for PD- L1(+) advanced gastric cancer patients as third-line treatment. In cohort 2, the enrollees were HER2(−) naïve patients with advanced gastric/gastroesophageal junction adenocarcinoma, who receive pembrolizumab associated with combined chemotherapy (5-fluorouracil/capecitabine plus cisplatin) for six cycles, and subsequently maintenance treatment with pembrolizumab plus 5-FU/capecitabine for a period of maximum 2 years/ until disease progression[228]. The preliminary data showed that 94% of patients have experienced adverse events (e.g., anorexia, nausea, neutropenia); of these patients, two thirds developed grade 3–4 toxicities. The results of the study proved manageable adverse events profiles for patients receiving combined therapeutic schemes, with none of the patients discontinuing the treatment. The data showed an ORR was 60% and 68.8% in PD-L1+ patients[229].

The KEYNOTE-028 phase Ib study evaluated the role of pembrolizumab administered in up to 2 years or until progression, in PD-L1(+) advanced solid tumors, including esophageal and gastroesophageal junction cancers (adenocarcinoma and squamous cell cancer), with most of them having at least 2 prior lines of chemotherapy[230]. The interim data on 23 patients showed an ORR of 30%, with 6- and 12-mo PFS rates of 30.4% and 21.7%, respectively. The response rates were better for adenocarcinomas.

An ongoing phase II study is assessing the efficiency of pembrolizumab monotherapy, along with the analysis of immune-related gene profiles and PD-L1 expression as biomarkers for treatment response in patients with advanced cancer of the esophagus or gastroesophageal junction (adenocarcinoma or squamous cell carcinoma) (KEYNOTE-180, NCT02559687)[231], progressing on standard chemotherapy.

KEYNOTE-061 is an ongoing phase III open-label clinical trial comparing pembrolizumab versus paclitaxel in the second-line setting for patients with advanced gastric or gastroesophageal cancer (progression after first-line therapy with a platinum + fluoropyrimidine combination) ([NCT02370498](http://clinicaltrials.gov/show/NCT02370498))[232]. The treatment administration continues until the disease progression or the occurrence of severe toxicities; the primary endpoints are PFS and OS in the PD-L1 positive population. The study randomized 592 patients to receive pembrolizumab or standard-dose paclitaxel. PD-L1 positivity was encountered in 395/592 patients enrolled. Median OS was 9.1 mo with pembrolizumab *vs* 8.3 mo with paclitaxel (HR = 0.82, *p* = 0.042); 12-mo OS rates in pembrolizumab group compared with paclitaxel group were 39.8% *vs* 27.1%, and 18-mo rates were 25.7% *vs* 14.8%. There was no difference in PFS or ORR, but pembrolizumab responses proved more durable, and the treatment effect was more prominent in patients with ECOG PS 0 (HR = 0.69), gastroesophageal junction tumors (HR 0.61) and with increasing PD-L1 expression. The safety profile observed in KEYNOTE-061 was consistent with that observed in previously reported studies of pembrolizumab, this drug demonstrating a better safety profile than paclitaxel. The agent reduced the risk of death by 18% *vs.* paclitaxel in patients with previously treated gastric cancer and PD-L1 positive, although this difference did not achieve statistical significance[233].

Furthermore, the ongoing phase III KEYNOTE-062 study is randomizing PD-L1(+)/HER2(-) advanced, metastatic gastric/GEJ adenocarcinoma patients to receive as a first-line regimen either pembrolizumab or pembrolizumab, associated with fluorouracil and cisplatin (NCT02494583)[234]. The primary endpoints are represented by OS and PFS.

A phase III study (KEYNOTE-181) aims to include approximately 600 patients with previously treated advanced adenocarcinoma or squamous cell carcinoma of the esophagus or gastroesophageal junction to receive either pembrolizumab or monochemotherapy (paclitaxel, docetaxel, or irinotecan) (NCT02564263)[235].

Anti-PD-L1 antibodies - Avelumab

Avelumab (MSB0010718C, trade name Bavencio) represents a fully human anti-PD-L1 IgG1 antibody which is approved by the FDA for the treatment of metastatic Merkel cell carcinoma (patients older than 12 years) and locally advanced/metastatic urothelial carcinoma under progression, after platinum-containing treatment (2017).

Avelumab is currently assessed in JAVELIN program (NCT01772004)[236]. Patients received 10 mg/kg of avelumab every 2 wk as second-line treatment. For Japanese gastric cancer patients, the obtained ORR was 15%, with 43.3% of patients presenting PFS at 12 wk[237]. In this context, phase Ib trials are currently investigating avelumab in patients with advanced gastric/gastroesophageal junction cancers. The results, presented at the American Society of Clinical Oncology (ASCO) 2016 meeting, showed that the most encountered adverse events were infusion-related reactions and fatigue. In the first context (patients who had at least one prior therapy), the ORR, DCR, and median PFS were 9.7%, 29.0%, and 6.0 wk, respectively, whereas in the second setting (patients who received avelumab as first line switch maintenance after chemotherapy), they were 9%, 57.3%, and 12.0 wk, respectively. ORR in the PD-L1- positive tumors was higher *vs.* PD-L1-negative tumors (18.2% *vs* 9.1% for the third-line treatment group and 10% *vs* 3.1% for maintenance group). Starting from these data, there are two phase III ongoing studies assessing avelumab in the treatment of gastric cancer patients. One study is comparing avelumab as third line of treatment to best supportive care (phase III JAVELIN Gastric 300 trial) (NCT02625623), while another is assessing patients receiving avelumab as a switch maintenance after chemotherapy with 5-FU or capecitabine plus oxaliplatin (phase III JAVELIN Gastric 100 study) (NCT02625610)[238]. The Javelin Gastric 300 trial enrolled 371 patients from 147 sites in Asia, Australia, Europe, North America and South America. Recently, the companies stated that the pivotal phase III Javelin trial investigating avelumab as third-line treatment for patients with unresectable, recurrent or metastatic gastric cancer did not meet its pre-specified primary endpoint of superior OS *vs* chemotherapy. It represents the first trial of a checkpoint inhibitor versus an active chemotherapy comparator rather than placebo in this hard-to-treat patient population. Safety profile was consistent with that observed in previously reported studies of avelumab. The JAVELIN Gastric 300 data will be further evaluated in an effort to better understand the results.

Anti-PD-L1 antibodies - Durvalumab (MEDI4736)

Durvalumab represents an engineered human anti-PD-L1 IgG1 antibody that prevents PD-L1 binding to PD-1 and CD80. It has received FDA approval for previously treated advanced bladder carcinoma and unresectable stage III NSCLC.

A phase I study assessed the efficiency of administering durvalumab in doses up to 10 mg/kg intravenously every 2 wk for up to one year in patients with solid tumors, including gastric cancer (NCT01693562)[239]. Fatigue, nausea, vomiting, rash and pyrexia were the most frequent adverse events. An ORR of 25% was obtained in patients with gastric tumors. Additionally, two patients with heavily pretreated tumors surpassed the current median PFS that was obtained with standard treatments.

An ongoing phase IB/II study is investigating patients with recurrent/metastatic gastric/gastroesophageal junction adenocarcinomas for monotherapy with durvalumab, tremelimumab, or a combination of durvalumab and tremelimumab (anti–CTLA-4) in second or third-line setting[240].

An algorithm of current treatment in unresectable locally advanced, recurrent or metastatic gastric cancer is presented in Figure 3.

**Checkpoint inhibitors in the adjuvant /neoadjuvant setting (stage II or III tumors):** Because multiple lines of standard chemotherapy may harm the immune system and immune biomarkers may be stimulated by previous chemoradiation, it was suggested that addition of checkpoint inhibition in the adjuvant or even neoadjuvant setting in earlier stages of the disease may induce a significantly higher tumor response rate[241]. It has been demonstrated that after chemoradiation neoadjuvant therapy, there is a higher number of TILs that are agglomerated in perivascular areas and arranged in lymphoid-like structures; these are features that favor the tumor response to immunotherapy[58].

A randomized phase III study (CheckMate-577) NCT02743494 is assessing the role of adjuvant nivolumab for patients with resected esophageal/gastroesophageal cancer. Investigators aim to enroll approximately 760 patients to receive either nivolumab or placebo. The primary endpoints are OS and DFS[242]. Moreover, a phase I study is currently investigating nivolumab combined with an anti-CCR4 (mogamulizumab) in the preoperative setting (NCT02946671)[243].

The combination of nivolumab and ipilimumab is going to be assessed in the adjuvant treatment of gastric/gastroesophageal junction adenocarcinomas with a high risk of recurrence – a phase II study (NCT03443856) (VESTIGE)[244].

Several ongoing phase I/II clinical trials are assessing the safety and efficacy of neoadjuvant treatment using different PD-1/PD-L1 checkpoint inhibitors,administered either concomitantly or sequentially with neoadjuvant chemoradiation. For example, a phase I trial is assessing neoadjuvant administration of nivolumab and ipilimumab in stage II/III patients (NCT03044613)[245]; another Ib phase clinical trial is assessing pembrolizumab and chemoradiation in the neoadjuvant setting of locally advanced esogastric tumors (NCT03064490/PROCEED)[246]. Several II and III phase ongoing studies are investigating the efficacy of pembrolizumab in association with chemotherapy as neoadjuvant/adjuvant treatment in gastric cancer patients[247-249]. Additionally, a phase II study is evaluating the results of administering avelumab plus chemotherapy (FLOT) in the perioperative setting[250].

**Future perspectives**

The synergistic action of the PD-1/PD-L1 checkpoint inhibitors and the vascular endothelial growth factor (VEGF)/vascular endothelial growth factor receptor (VEGFR) blockade was demonstrated by preclinical data[251]; the combined treatment induced an improved tumor response and better OS. This combination treatment assessed in phase I/II studies was associated with an acceptable safety profile [252,253].

A phase Ia/Ib study (NCT02443324)[254] is investigating the safety and efficacy of combining anti-PD-L1 (durvalumab) and anti-VEGFR2 antibodies (ramucirumab) in patients with refractory gastric/gastroesophageal junction tumors. The interim data of 40 patients showed a 45% DCR and a median PFS of 2.60 mo. The most frequently encountered toxicities included fatigue, infusion-related reaction, loss of appetite, pruritus, rash, and hypertension; 25% of patients had severe toxicities.

Additionally, the efficacy of associating nivolumab with ramucirumab in patients with advanced, unresectable gastric/gastroesophageal junction cancers is currently under investigation in a phase I/II study (NCT02999295)[255]. The clinical trial of nivolumab with paclitaxel and ramucirumab in patients with advanced, unresectable gastric cancer is ongoing under investigation in a phase I/II study (UMIN000025947)[256].

A phase II study is currently recruiting patients with metastatic/recurrent gastric/gastroesophageal cancer to receive pembrolizumab plus lenvatinib mesylate (anti-VEGFR2 tyrosine kinase inhibitor) (NCT03413397)[257], whereas another phase I/II study is planning to combine avelumab with regorafenib (another anti-VEGFR2 tyrosine kinase inhibitor) in advanced gastric tumors (NCT03475953/REGOMUNE)[258].

Because HER2+ tumors develop resistance to trastuzumab, studies have been evaluating a combination of PD-1 blockade plus anti-HER2 agents. Preclinical studies indicate that the Her2 blockade stimulates T cell activation and enhances interferon-γ secretion by NK cells and antibody-dependent cellular toxicity; therefore, it boosts the PD-1/L1 inhibitors’ efficacy[259].

The combination of pembrolizumab plus anti-HER2 monoclonal antibody margetuximab in patients with advanced HER2+ gastric cancers that are resistant to classical trastuzumab-based chemotherapy is currently under evaluation in a phase Ib/II dose-escalation trial (NCT02689284)[260].

Other promising options include dual immunotherapies, such as the administration of PD-1/PD-L1 checkpoint inhibitors combined with agents suppressing other immune checkpoints (*e.g.*, TIM3, LAG3) or T cell costimulatory antibodies (*e.g.*, GITR, OX40, and 4-1BB). Additionally, other directions investigate the combination of the PD-1/PD-L1 blockade with radiation and other cytotoxic and targeted treatments (PARP inhibitors, ATR serine/threonine protein kinase inhibitors, pegylated recombinant human hyaluronidase, anti-CEACAM1, arginase inhibitor INCB001158, FGFR inhibitors, immunotherapy using Listeria bacteria to activate immune response against specific tumor-associated antigens, claudiximab, *etc*.), and enzymatic inhibitors such as IDO-1[211].

Therapeutic intervention upon IDO1-mediated immune suppression involves inhibition of the catalytic activity of IDO1. Preliminary data from the ongoing studies investigating dual inhibition of both PD-1 and IDO1 in solid tumors are promising, showing an increased efficiency of the checkpoint blockade[261,262]. Currently, there are two studies of phase I/II (NCT02178722/KEYNOTE-037/ECHO-202)[263] and II (NCT03196232)[264], respectively investigating the IDO1 inhibitor epacadostat in combination with pembrolizumab in gastric tumors.

Additionally, CDC20 encoding cell-division cycle protein 20 homologue[265], as well as PLK1 and TTK, which represent checkpoints in mitosis, may be used as therapeutic targets[266]. PLK1 is upregulated in tumors, and data reported from both preclinical and clinical studies suggest encouraging benefits from PLK-1 inhibition[267]. Maternal embryonic leucine zipper kinase (MELK) decreases the apoptosis of tumor cells; therefore, additional MELK inhibition may also be offered in PD-L1+ cancers[268].

The negative checkpoint regulator VISTA represents a V-domain Ig suppressor of T-cell activation (VISTA), known as PD1 homolog, which belongs to the B7 family. VISTA is mostly exhibited on hematopoietic cells[269]. In murine tumor models, anti-VISTA monoclonal antibodies activated intratumoral T cells, enhancing antitumor immunity. Moreover, the combined VISTA/PD-1 blockade was proved efficient[270,271]. A phase I study, combining VISTA and PD-L1/PD-L2 inhibition in solid tumors (CA-170 molecule; NCT02812875), has been initiated[272]. Böger et al., investigated VISTA expression in 464 gastric cancer patients, with an increased expression associated with the intestinal type (VISTA is a regulator of differentiation), proximal gastric tumors and KRAS (-) and PIK3CA mutant cancers. Additionally, a significantly higher VISTA expression in immune cells was noticed in association with PD-L1 expression and in EBV+ tumors. These observations might stress that a subset of tumors may use multiple checkpoint pathways to escape immunity and that the combined VISTA/PD-1 inhibition may be a promising novel cancer approach[273].

Another study is investigating nivolumab plus a matrix metalloproteinase 9 inhibitor (GS-5745 = andecaliximab) in patients with unresectable/recurrent gastric or gastroesophageal junction adenocarcinoma (NCT02864381)[274].

Due to the permanent development of novel immunotherapeutic agents, classical trials do not have the ability to appropriately assess all possible drug combinations. In this regard, FRACTION (Fast Real-time Assessment of Combination Therapies in Immuno-Oncology) is a new clinical trial program with a dynamic design that provides the possibility both for the inclusion of new immunotherapeutic combinations, as well as exclusion of ineffective ones[275].

Cancer radiotherapy may seldom exhibit the clinical phenomenon of the abscopal response, meaning that no irradiated metastases diminish after radiation to the primary tumor. Data from preclinical trials highlighted that radiotherapy plus PD-1/PD-L1 blockade exhibited synergistic anticancer effects. Currently, trials on gastric cancer patients are investigating the efficiency of pembrolizumab plus palliative radiotherapy (metastatic tumors), and additional neoadjuvant chemoradiotherapy (resectable tumors) (NCT02730546)[276], respectively. A phase I/II trial is also investigating nivolumab + radiotherapy in unresectable, recurrent gastric cancer (as third-line treatment) (NCT03453164) (CIRCUIT)[277].

**Predictive biomarkers and genetic profiles**

The combination of immune checkpoint inhibitors with other types of immunotherapies, targeted agents, radio- or chemo-therapy, seems to generate promising results against gastrointestinal cancers but at the expense of the occurrence of immune-related side effects, some of them potentially fatal[278,279]. Therefore, identification of prognostic biomarkers and genetic profiles to define subgroups of gastric cancer patients who are most likely to respond to specific immunotherapeutic regimens is an urgent need[280].

In this regard, an ongoing observational study NCT02951091 (Biomarker – integrated Umbrella)[281] is investigating different molecular cohorts in oncologic patients, the cases of PD-L1+, MSI-H, and EBV+ advanced gastric cancers being assigned to receive either nivolumab or other agents, such as AFATINIB (EGFR tyrosine kinase inhibitor or GSK2636771 (PI3K beta inhibitor) +PACLITAXEL). A phase II study compares nivolumab with other novel agents, according to genetic testing, in gastric cancer patients with mismatch repair deficiency (loss of MLH1/ MLH2) (NCT02465060 - The MATCH screening trial)[282].

**Conclusion**

because of the well-known heterogeneity of tumors, it is essential to assess the particular molecular biology of different subtypes of gastric cancers that are associated with different clinico-biologic parameters and prognosis to identify innovative treatment approaches that will improve current results in gastric cancer. Modern treatments, such as the promising immunotherapy, should be applied in this way to selected patients who have been proven to have the best response to a specific therapy (individualized treatment). In this context, it is also mandatory to discover prognostic tumor markers and predictive biomarkers for treatment response.

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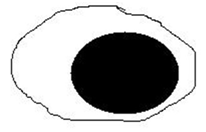
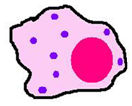
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**A**



NK

Th CD4+

Tc CD8+

**IFN γ**

inhibits angiogenesis proliferation

P

phagocytosis

**+**

**+**

**-**

CC

angiogenesis

DC

Macrophage

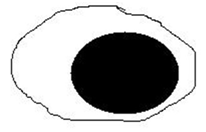
**Elimination phase**

**B**

**Equilibrium phase**

**IFN γ**

**IL 12**



Tc CD8+

DC

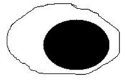
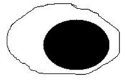
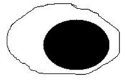
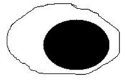
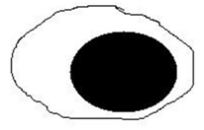
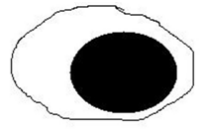
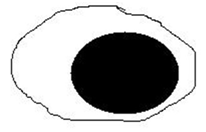
CC

**-**

angiogenesis

**C**

**Escape phase**



Clonal expansion

CC

PD-L1

PD-1

**-**

Tc CD8+

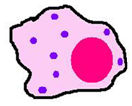
Th CD4+

NK

T reg.

**-**

Blood vessels



Macrophage

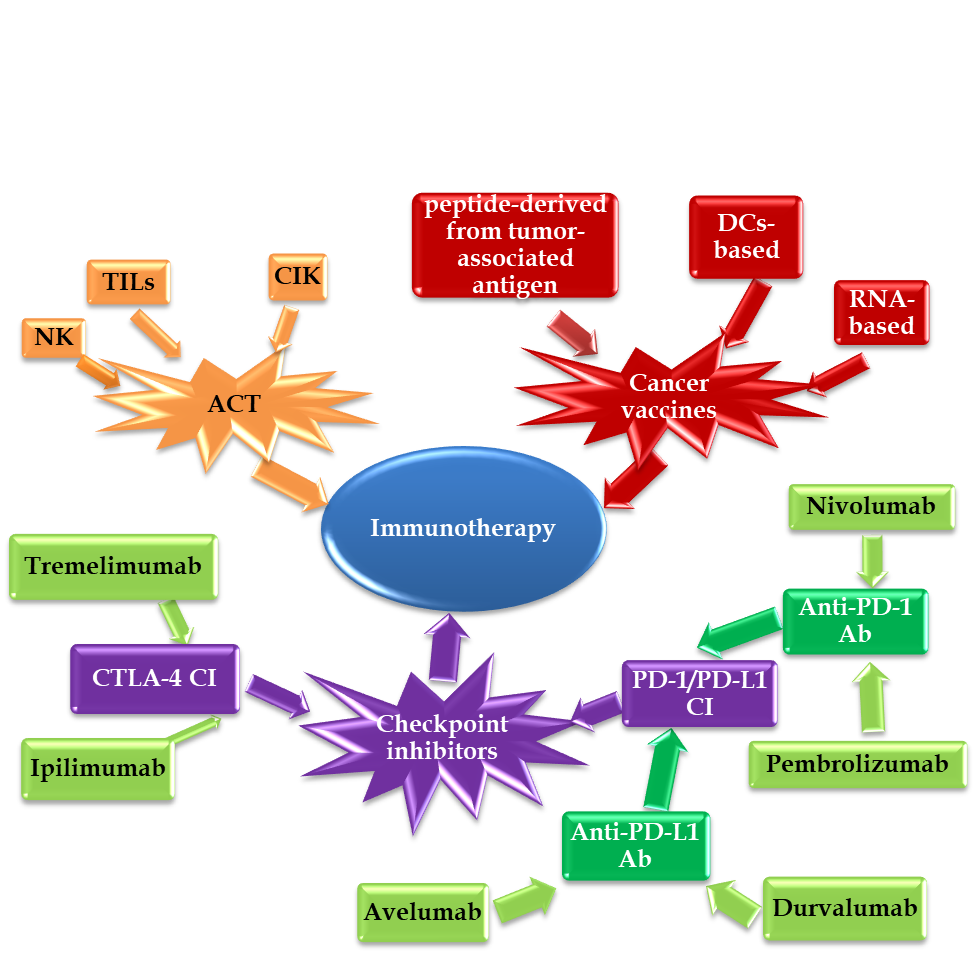
TGF β

**-**

**+**

Neoangiogenesis proliferation

**Figure 1 The three phases of cancer immunoediting.** A: Phase 1: Elimination; B: Phase 2: Equilibrium; C: Phase 3: Escape.



**Figure 2 Current immunotherapeutic strategies in gastric cancer.** NK: natural killer cells; TILs: tumor-infiltrating lymphocytes; CIK: cytokine-induced killer cells; ACT: adoptive cell therapy; DCs: dendritic cells; PD-1: programmed death-1; PD-L1: programmed death ligand-1; CTLA-4: cytotoxic T-lymphocyte-associated antigen 4; CI: checkpoint inhibitors.

**I st line treatment**

**4 th line**

**PD-L1 + tumors**

**3rd line**

**2nd line**

**HER 2 -**

**HER 2+**

**Figure 3 Therapeutic algorithm in unresectable locally advanced, recurrent or metastatic gastric cancer.** HER2: human epidermal growth factor receptor 2; CX: cisplatin and capecitabine; CF: cisplatin and fluorouracil; MSI: microsatellite instability; dMMR: deficient mismatch repair gene; PD-L1: programmed death ligand-1.

**Table 1 Clinical trials using CTLA-4 and PD/PD-L1 checkpoint inhibitors**

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Checkpoint inhibitors | Study number | Phase | Status | Study design | Study population | Primary outcome measures | Secondary outcome measures | No. of patients (estimated enrollment) | Final results |
| Nivolumab, Ipilimumab | NCT03342417 | II | recruiting | NIVO + IPI | -neoadjuvant breast cancer  -platinum-resistant advanced ovarian/GC | - % AE | - DOR, OS, QOL, recurrence rate | 60 | pending |
| Nivolumab, Ipilimumab | NCT02872116  (CHECKMATE-649) | III | recruiting | - NIVO + IPI/  -NIVO + chemo  vs. chemotherapy (XELOX/FOLFOX) | -naive advanced/metastatic GC/GEJ | -OS NIVO+IPI vs. chemo (PD-L1+ tumors)  -OS NIVO+chemo vs. chemo  -ORR, PFS nivo+chemo vs. chemo | -OS NIVO+IPI vs. chemo  -PFS NIVO+IPI/  NIVO+chemo vs. chemo (PD-L1+)  -ORR NIVO+chemo vs. chemo | 1349 | pending |
| Nivolumab, Ipilimumab | NCT02935634  (FRACTION-GC) | II | recruiting | - NIVO + IPI  -NIVO+RELATLIMAB  -NIVO+BMS-986205 | -advanced GC | -ORR, DOR, PFS | -% AE, SAE, discontinuation/death due to treatment | 300 | pending |
| Nivolumab, Ipilimumab | NCT03044613 | Ib | recruiting | - NIVO/NIVO + IPI prior to chemoradiation+NIVO | -neoadjuvant treatment, resectable stage II/III EC/GEJ | -% AE | -feasibility of induction treatment  -path CR  -quantity of NIVO bound to PD1 receptor  -changes in expression of immune markers  -OS, RFS | 32 | pending |
| Nivolumab, Ipilimumab | NCT03443856  (VESTIGE) | II | Not yet recruiting | - NIVO + IPI | -adjuvant treatment GC/GEJ adenocarcinoma stage Ib-IVa, ↑risk of recurrence (ypN1-3+/R1) after neoadjuvant treatment +resection | -DFS | -OS  -relapse rate  -loco-regional/distant failure rates  -% AE, QOL, global health status | 240 | N/A |
| Nivolumab, Ipilimumab | NCT03409848  (INTEGA) | II | recruiting | -IPI/FOLFOX+ NIVO & TRAS | -advanced/metastatic GC adenocarcinoma, previously untreated | -OS | -% AE  -PFS, RR, QOL  -translational research  -central imaging review | 97 | pending |
| Nivolumab, Ipilimumab | NCT02834013 | II | recruiting | -NIVO+IPI | -rare tumors, including gastric NET, SqCC, GIST | -ORR-RECIST | -toxicities  -OS, PFS  -clinical benefit rate  -ir-ORR/PFS | 707 | pending |
| Nivolumab, Ipilimumab | NCT03126110 | I/II | recruiting | -NIVO+INCAGN01876  -IPI+ INCAGN01876  -NIVO+IPI+ INCAGN01876 | -advanced/metastatic malignancies, including GC | -toxicities (%AE)  -ORR-RECIST | -DR, DDC, PFS, OS- RECIST | 450 | pending |
| Nivolumab, Ipilimumab | NCT03241173 | I/II | recruiting | -NIVO+INCAGN0949  -IPI+ INCAGN0949  -NIVO+IPI+ INCAGN0949 | -advanced/ metastatic malignancies, including GC | -toxicities (%AE)  -ORR-RECIST | -DR, DDC, PFS, OS- RECIST | 651 | pending |
| Ipilimumab | NCT01585987 | II | completed | -IPI vs. BSC (5-FU) | -unresectable/ metastatic GC/GEJ adenocarcinoma (following Ist line treatment) | -ir-PFS (ir RECIST)  -% BOR | -PFS (mWHO)  -OS | 143 | -ir-PFS ↓ (2.92  vs. 4.89  months);  -mWHO-PFS↓  (2.72 vs. 4.89)  (p=0.03);  -OS at study completion  12.68 vs.  12.06 months |
| Tremelimumab, Durvalumab | NCT02658214 | I | recruiting | -TREME+DURVA+chemo (platinum-based SOC) | -Ist line locally advanced/ metastatic solid tumor, including GC/GEJ | -laboratory findings  -%AE, safety, tolerability  -tumor assessment (RECIST) | - | 42 | pending |
| Nivolumab | NCT03453164 (CIRCUIT) | I/II | recruiting | NIVO+radiotherapy | -unresectable recurrent GC (3rd line) | -DCR:  -PD: on CT/ MRI/ PET-CT  -CR/PR/SD | -mean survival  -% AE  -local control rate  -% PL-L1+, MHCI- tumor cells  -cytokines serum concentration  -% regT cells  -% Ag-specific CTL | 40 | pending |
| Nivolumab | NCT02267343 | III | active, not recruiting | NIVO vs. placebo | -refractory, unresectable, advanced/ recurrent GC/GEJ | -OS, PFS | -ORR, DOR, %AE, %SAE, safety | 480 | pending |
| Nivolumab | NCT02746796 | II/III | recruiting | NIVO+chemo | -Ist line therapy, unresectable advanced/ recurrent GC/GEJ | -PFS  -OS | -ORR, DOR, DCR  -TTR, BOR  -% AE, SAE, laboratory abnormalities | 680 | pending |
| Nivolumab | NCT03006705 | III | recruiting | NIVO+chemo vs. placebo+chemo | -adjuvant treatment p stage III GC/GEJ (after D2 resection) | -RFS | -OS  -safety- % AE, SAE, laboratory abnormalities | 700 | pending |
| Nivolumab | NCT02999295 | I/II | recruiting | -NIVO+RAMUCIRUMAB | -advanced/ recurrent unresectable GC/GEJ | -no. of pts with DLT  -6 mo PFS | -%AE  -ORR  -DCR  -OS, PFS | 44 | pending |
| Nivolumab | NCT02946671 | I | recruiting | -NIVO+MOGAMULIMUMAB (KW-0761 = anti-CCR4) | -preoperator treatment against solid cancers, including GC | -% AE  - FOXp3+ tumors by immunohistochemistry | -ORR-RECIST  -% ↓Treg | 18 | pending |
| Nivolumab | NCT02951091 (Biomarker – integrated Umbrella) | observational | recruiting | -NIVO/ AFATINIB/ GSK2636771+PACLITAXEL | -advanced GC  -different molecular cohorts: PD-L1+, MSI-H, EBV+ → NIVO | -PFS | - | 400 | pending |
| Nivolumab | NCT02465060 (The MATCH screening trial) | II | recruiting | -NIVO/ other agents (according to genetic testing) | -advanced/ metastatic solid tumors (including GC), lymphomas, multiple myelomas → mismatch repair deficiency (loss of MLH1/ MLH2) | -ORR | -OS, PFS  -TTP | 6452 | pending |
| Nivolumab | NCT02862535 | Ib | active, non-recruiting | -ANDECALIXIMAB (GS-5745) ± NIVO/chemo | -previously treated, advanced GC/GEJ adenocarcinoma (Japan) | - safety (% AE) | - serum concentration of Andecaliximab  -% Ab-anti Andecaliximab | 36 | N/A |
| Pembrolizumab | NCT03382600 (MK-3475-659/ KEYNOTE 659) | II | recruiting | -PEMBRO+OXALIPLATIN+TS-1 vs.  -PEMBRO+CISPLATIN+TS-1 | -advanced CG/GEJ adenocarcinoma, HER2(-), PD-L1+ | -ORR-RECIST | -ORR (i-RECIST)  -DCR, DOR, TTR, PFS (RECIST, iRECIST)  -OS, % AE | 90 | pending |
| Pembrolizumab | NCT02901301 | I/II | recruiting | -PEMBRO + TRASTUZUMAB + chemo | -Ist line advanced, GC HER2+ | -recommended dose  -ORR-RECIST | -DOR  -TTR | 49 | pending |
| Pembrolizumab | NCT03342937 | II | recruiting | PEMBRO+XELOX | -Ist line metastatic GC adenocarcinoma | -PFS | -OS  - % AE | 50 | pending |
| Pembrolizumab | NCT02918161 | II | recruiting | PEMBRO+chemo (SOC) | -perioperative setting GC/GEJ | -2 years DFS | - pathol CR  -OS, ORR (RECIST)  -DFS | 40 | pending |
| Pembrolizumab | NCT02689284 | I/II | recruiting | PEMBRO+MARGETUXIMAB | -HER2+ advanced, metastatic GC/GEJ | - expansion phase dose of Margetuximab  -antitumor activity: RD, ORR (RECIST, ir-RECIST) | -OS  -PFS | 72 | pending |
| Pembrolizumab | NCT03257163 | II | recruiting | -PEMBRO+CAPECITABINE+radiotherapy (perioperative) | -mismatch repair deficient, EBV+, operable GC | -RFS | - | 40 | pending |
| Pembrolizumab | NCT03064490 (PROCEED) | II | recruiting | -PEMBRO+chemoradiotherapy | -neoadjuvant treatment, locally advanced EG cancers | -pathol CR | -toxicity (% AE) | 38 | pending |
| Pembrolizumab | NCT02563548 | Ib | recruiting | -PEMBRO+PEGPH2O (Pegylated Recombinant Human Hyaluronidase) | -hyaluronan-high (HA-H) patients with relapsed/ refractory cancers (adenocarcinoma) | -ORR | -DCR, DOR, PFS (RECIST, ir-RECIST) | 81 | pending |
| Pembrolizumab | NCT02954536 | II | recruiting | -PEMBRO+TRASTUZUMAB+chemo | -advanced, metastatic HER2+, EG (Ist line) | -PFS (RECIST) | - | 37 | pending |
| Pembrolizumab | NCT03221426 (MK-3475-585) (KEYNOTE-585) | III | recruiting | -PEMBRO+cemo vs. placebo+chemo | -neoadjuvant/adjuvant previously untreated GC/GEJ adenocarcinoma | -OS  - EFS event-free survival)  -Pathol CR  -% AE + discontinuation of treatment | -DFS | 860 | pending |
| Pembrolizumab | NCT03196232 | II | recruiting | -PEMBRO+EPACADOSTAT | -metastatic/ unresectable GEJ | - 6-mo PFS | -ORR (RECIST)  -OS  -RR | 30 | pending |
| Pembrolizumab | NCT03019588 (MK-3475-063/ KEYNOTE-063) | III | recruiting | -PEMBRO vs. chemo (PACLITAXEL) | -progression after Ist line platinum-fluoropyridine chemo, advanced GC/GEJ adenocarcinoma, PD-L1+ (Asia) | -OS, PFS (RECIST) | -ORR-RECIST  -% AE, % discontinuation due to AE | 360 | pending |
| Pembrolizumab | NCT03488667 | II | not yet recruiting | PEMBRO+mFOLFOX | -neoadjuvant treatment GEJ adenocarcinoma  -adjuvant treatment GC | -yp RR (pathologic response)  -toxicity (% AE) | -ORR, DFS, OS  -PET scan response rate  - % PD-L1+ in tumor cells | 40 | N/A |
| Pembrolizumab | NCT03413397 | II | recruiting | PEMBRO+LENVATINIB MESYLATE | -metastatic/ recurrent GC/GEJ | -ORR-RECIST | -PFS, OS  -characteristic immunologic changes | 29 | pending |
| Pembrolizumab | NCT02730546 | I/II | recruiting | PEMBRO+chemoradiotherapy | -locally advanced, operable, GEJ/gastric cardia adenocarcinoma (neoadjuvant setting) | -path CR  -PFS | -R0 resection, DSF  -dose-limiting AE, % surgical complications, OS, PFS, time to relapse | 68 | pending |
| Pembrolizumab | NCT02318901 | Ib/II | active, not recruiting | PEMBRO- TRASTUZUMAB/ ADO-TRASTUZUMAB-ETAMSINE/ CETUXIMAB | -unresectable HER2+, advanced GC/GEJ | -dose of mAb combined with PEMBRO | -% grade 3-4 AE  -RR (RECIST, ir-RECIST)  -OS, PFS  -circulating tumor DNA  -imaging changes | 90 | N/A |
| Pembrolizumab | NCT03095781 | I | recruiting | PEMBRO+XL888 (= Hsp90 inhibitor) | -advanced gastrointestinal cancer (including GC) | -recommended dose for combined treatment | -ORR, PFS, RS (RECIST)  -OS | 50 | pending |
| Pembrolizumab | NCT02346955 | I | terminated | -CM-24[MK-6018 = mAb against CEACAM1] ± PEMBRO | -advanced/recurrent malignancies (including GC) | -% AE, discontinuation due to AE, DLT | -maximum drug concentration, half-life elimination, ORR, DOR | 27 | pending |
| Pembrolizumab | NCT02178722 (KEYNOTE-037/ECHO-202) | I/II | recruiting | -PEMBRO+EPACADOSTAT | -selected carcinomas (including GC) | -% DTL  -ORR | -PFS  -% AE  -OS | 508 | pending |
| Pembrolizumab | NCT02903914 | I/II | recruiting | -PEMBRO+ARGINASE INHIBITOR INCB001158 | - advanced/ metastatic solid tumors (including GC) | -% AE | - recommended dose of arginase inhibitor ± PEMBRO  - pharmacokinetic profile  -antitumor activity of drugs (RECIST, ir RECIST) | 346 | pending |
| Pembrolizumab | NCT03122548 | II | Active, not recruiting | PEMBRO+CRS-207 | -recurrent/ metastatic GC/EG (1-2 prior lines of systemic treatment) | -% AE | -tumor response (RECIST)  -OS  -characterization of immune response  -analysis of biomarker expression | 79 | N/A |
| Pembrolizumab | NCT02393248 | I/II | recruiting | -INCB054828+PEMBRO/ chemo/ TRASTUZUMAB | -advanced malignancies (including GC), progression after prior treatment | -maximum tolerated dose, pharmacodynamic of INCB054828 | -ORR  -maximum/minimum plasma concentration of NCB054828 | 280 | pending |
| Pembrolizumab | NCT02494583 | III | active, not recruiting | -PEMBRO vs.  -PEMBRO+chemo vs.  -placebo+chemo | -Ist line treatment, advanced GC/GEJ | -PFS (RECIST)  -OS | -ORR, DOR (RECIST)  -QOL | 764 | N/A |
| Pembrolizumab | NCT02370498 (KEYNOTE-061) | III | active, not recruiting | -PEMBRO vs. chemo (PACLITAXEL) | -advanced GC/GEJ adenocarcinoma, progressed after Ist line (platinum+fluoropyrimidine), PD-L1+ | -PFS, OS- in PD-L1+ | -PFS, OS  -TTP, ORR | 592 | pending |
| Pembrolizumab | NCT02335411 (KEYNOTE-059) | II | active, not recruiting | -PEMBRO or PEMBRO+chemo (CISPLATIN+5-FU/CAPECITABINE) | -recurrent/metastatic GC/GEJ adenocarcinoma | -% AE, discontinuation of treatment due to AE  -ORR | - | 316 | pending |
| Pembrolizumab | NCT03277352 | I/II | recruiting | -INCAGN01876+PEMBRO+EPACADOSTAT | -advanced/ metastatic malignancies | -% AE  -ORR, CRR (RECIST) | -ORR, DCR, DOR, PFS, OS (rECIST, mRECIST) | 166 | pending |
| Pembrolizumab | NCT02443324 | I | active, not recruiting | -PEMBRO+RAMUCIRUMAB | -GC/GEJ (NSCLC, transitional urothelial cancer, biliary tract cancer) | -DTL | -% BOR of CR/PR, ORR  -% SD  -DOR, time to response  -PFS, OS  -pharmacokinetics | 155 | pending |
| Avelumab | NCT02625623 (JAVELIN GASTRIC 300) | III | active, not recruiting | -AVE+ BSC vs.  -chemo+BSC/BSC | -unresectable, recurrent, locally advanced/ metastatic GC/GEJ adenocarcinoma (3rd line) | -OS | -PFS, BOR  -QOL | 37 | pending |
| Avelumab | NCT03399071 (ICONIC) | II | recruiting | -AVE+chemo (FLOT) | -perioperative setting, operable EC/GC | -pathol CR | -% grade 3-4 AE  -radiologic response (RECIST)  -median PFS, OS | 40 | pending |
| Avelumab | NCT02625610 (JAVELIN GASTRIC 100) | III | active, not recruiting | -AVE maintenance vs. Ist line continuation of chemo (OXALIPLATIN+FLUOROPYRIMIDINE) | -unresectable, locally advanced/ metastatic GC/GEJ adenocarcinoma | -OS, PFS | -BOR (RECIST)  -QOL  -% AE | 499 | pending |
| Avelumab | NCT01943461 (JAVELIN SOLID TUMOR JPN) | I | active, not recruiting | -AVE | -locally advanced/ metastatic solid tumors (Japan) → expansion part GC patients (Asia) | -DLT | -concentration assessment, elimination half-life  -% PD-L1  -BOR+ irBOR, PFS +irPFS  -OS  % Ab-anti AVE  -% AE | 57 | pending |
| Avelumab | NCT03475953 (REGOMUNE) | I/II | not yet recruiting | -AVE+REGORAFENIB | -advanced/ metastatic digestive solid tumors (including GC) | -recommended doses  -assessment of Regorafenib antitumor activity  -pharmacokinetics | -maximum tolerated dose  -DLT  -% AE  -BOR, ORR, PFS, OS  -blood/tumor growth biomarkers | 212 | pending |
| Avelumab | NCT02554812  (JAVELINE Medley) | Ib/II | recruiting | -AVE+other immunotherapies → AVE+PD 0360324 (M-CSF mAb) (gastric cancer) | -locally advanced/ metastatic solid tumors (including GC) | -% DLT  -ORR | -serum concentration of drugs  -Ab-anti drugs  -TTR, DOR, PFS, OS  -tumor biomarkers (PD-L1, CD8+T cells) | 560 | pending |
| Durvalumab | NCT02734004 (MEDIOLA) | I/II | active, not recruiting | -DURVA+OLAPARIB (PARP inhibitor) | -advanced solid tumors (including GC) | -DCR (at CT/MRI)  -% AE  -safety- vital signs, blood samples | -% PD-L1  -DCR (mRECIST)  -time to treatment discontinuation)  -OS  -% change in tumor size (CT/MRI)  -serum concentration of Ab-anti drug  -pharmacokinetics  -ORR, DOR, PFS (mRECIST) | 148 | pending |
| Durvalumab | NCT02572687 | I | active, not recruiting | DURVA+RAMUCIRUMAB | -locally advanced unresectable/ metastatic gastrointestinal (including GC/GEJ adenocarcinoma) and thoracic malignancies | -% DLT | -ORR, DCR  -DOR, TTR  -PFS  -OS  -pharmacokinetics  -% Ab-anti drug | 114 | pending |
| Durvalumab | NCT02678182 (PLATFORM) | II | recruiting | -DURVA vs. CAPECITABINE vs. TRASTUZUMAB vs. RUCAPARIB vs. surveillance | - maintenance treatment, locally advanced/ metastatic HER2+/- EG, adenocarcinoma (after Ist line chemo) | -PFS (RECIST) | -PFR  -OS, ORR (RECIST)  -% AE  -PFS, PFR, OS, ORR according to PD-L1 immunohistochemical status | 770 | pending |
| Durvalumab | NCT02264678 | I/II | recruiting | -AZD6738±Chemo/OLAPARIB/DURVA | -advanced malignancies (including GC) | -safety, tolerability (AE, SAE) | -pharmacodynamics, biomarker changes  -serum concentration of drug, half-life, etc.  -BOR, ORR, % change in tumor size | 250 | pending |

NIVO: nivolumab; IPI: ipilimumab; TREME: tremelimumab; DURVA: durvalumab; PEMBRO: pembrolizumab; AVE: avelumab, TRAS: trastuzumab; chemo: chemotherapy; AE: adverse events; SAE: serious adverse events; DLT: dose limiting toxicities; DOR: duration of response; OS: overall survival; QOL: quality of life; PFS: progression free disease; ir-PFS: immune-related progression free disease; PFR: progression-free rate; ORR: objective response rate; RR: response rate; pathCR: pathologic complete response, RFS: relapse-free survival; DR: disease response; DDC: duration disease control; BOR: best objective response; DCR: disease control rate; PD: progressive disease; CR: complete response; PR: partial response; SD: stable disease; TTR: time to response; NET: neuroendocrine tumors; SqCC: squamous cell carcinoma; GIST: gastrointestinal stromal tumors; EBV: Epstein Barr virus; N/A: not applicable.