# Anti-CTLA-4 therapy for malignant mesothelioma

Guazzelli, A, Bakker, EY, Krstic-Demonacos, M, Lisanti, MP, Sotgia, F and Mutti, L

http://dx.doi.org/10.2217/imt-2016-0123

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<td><strong>Published Date</strong></td>
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**Abstract**

Immunotherapy is an emerging therapeutic strategy with a promising clinical outcome in some solid tumours, particularly metastatic melanoma. One approach to immunotherapy is immune checkpoint inhibitors, such as blockage of cytotoxic T-lymphocyte antigen-4 (CTLA-4) and programmed death-1 (PD-1) and/or programmed death-ligand 1 (PD-L1).

This special report aims to describe the state of clinical trials of tremelimumab in patients with unresectable Malignant Mesothelioma (MM) in particular with regard to the clinical efficacy, safety and tolerability. Criticism and perspective of this treatment are also discussed.

Biological and clinical considerations rule out the use of tremelimumab as single agent for MM and, more in general, the use of immune checkpoint inhibitors for MM is still largely questionable and not supported by evidences.

**Keywords:** Malignant mesothelioma, immunotherapy, anti-CTLA-4
1. Introduction

Malignant Mesothelioma (MM) is an aggressive tumour of mesothelial cells that may be pleural (>80%), pericardial, peritoneal or vaginal/testicular and is associated with asbestos exposure [1, 2]. Although rare, MM incidence is increasing, especially in industrialized nations where asbestos is not banned [3] and the incidence is expected to peak between 2015-2030 [4]. Simian virus 40, radiation, erionite and genetic susceptibility may contribute to MM development [5, 6].

MM has poor prognosis (median survival of 9-12 months) and the majority of patients are diagnosed in an advanced stage [1]. The lack of effective diagnostic tools for early detection and a lack of validated circulating biomarkers complicate diagnosis [7-11]. Combination chemotherapy with antifolate/cisplatin is the standard first-line treatment, which prolongs median survival by just 2.8 months [12]. Second-line treatments are not still established. The need to discover new druggable targets is urgently required to overcome the poor prognosis of MM. Immunotherapy, in particular immune checkpoint blockade, is currently one of the most explored and attractive therapeutic approaches in oncology [13]. Immune checkpoint therapy, using blocking antibodies to CTLA-4 and PD-1/ PD-L1 aim to aid the immune system in the elimination of cancer cells [14]. In this special report we will discuss and comment on current clinical studies of anti-CTLA-4 therapy alone and in combination with PD1/PD-L1 inhibitors as double-checkpoint therapy in patients with MM.

2. Immunotherapy

Immunotherapy is an emerging therapy that has acquired clinical interest in cancer treatment [15]. The use of monoclonal antibodies (mAbs) in cancer has shown good promise. MAbs interfere with a single target molecule, with high selectivity due to their specificity [16]. Therefore, the innovative principle of immunotherapy is to not target cancer cells but “educate” the immune system to recognise neoplastic cells. Theoretically, immunotherapy could provide long-term disease control and improve survival by limiting tumour expansion [17].

Ipilimumab and tremelimumab (CP-675,206), both mAbs against CTLA-4, are the main clinically developed immunotherapy drugs and have entered advanced clinical trials for several cancers. Both are fully human novel mAbs, which are able to interrupt co-inhibitory pathways, promoting T-cell activation and tumour-specific cytotoxicity (Fig. 1) [18]. Ipilimumab is an immunoglobulin IgG1
isotype and has a half-life of 12 to 14 days. Tremelimumab is a non-complement-fixing IgG2 isotype, which causes less antibody-dependent cellular toxicity than IgG1 and has a half-life of 22 days, thus making tremelimumab a potentially better candidate than ipilimumab [19]. Ipilimumab has been approved by Food and Drug administration (FDA) in unresectable advanced stage of melanoma [20]. However, tremelimumab remains under investigation at different doses as monotherapy or in combinations in different tumours, including MM.

After the success of ipilimumab in cutaneous melanoma patients, additional checkpoint-blocking mAbs have been developed against PD-1 and its ligand CD274 (PD-L1). Nivolumab (Opdivo®, BMS-936558, MDX1106) and pembrolizumab (Keytruda® MK-3475, lambrolizumab) represent the first generation of humanized anti-PD-1 antibodies and currently are the main PD-1/PD-L1 blockade approved by the FDA to treat advanced melanoma [21].

3. Immunotherapy mechanisms of action and the rationale behind the preclinical studies

CTLA-4 (CD152) is a regulatory molecule, involved in the negative regulation of T-cells [22, 23]. CTLA-4 receptor and the co-stimulatory receptor CD28 share identical ligands: CD80 (B7.1) and CD86 (B7.2) and have the role of interfering the CD28-B7 axis, resulting in a reduction of T-cell proliferation and a decrease in interleukin-2 (IL-2) production [24]. Effector T-cells regulate CTLA-4 expression on the plasma membrane after induction of the co-stimulatory interaction between CD28 and B7 on APCs (Fig. 1). T_reg cells, however, express CTLA-4 constitutively to maintain their suppressive function [25]. In cancer, it has been shown that the predominance of co-inhibitory pathways influences the feature of tumour evasion from immune system control [26, 27]. Therefore, the discovery of this negative signalling receptor and its critical role in tumour-induced immune tolerance by in vitro studies and preclinical models has allowed the development of a new class of immunotherapeutic drugs [28-30]. Indeed, the crucial function of CTLA4 for negatively regulating T-cell activation has been demonstrated by the fatal systemic immune hyperactivation phenotype of Ctlb4-knockout mice, providing the rationale for anti-CTLA-4 therapy [28, 29].

Another study using mesothelioma-bearing mice showed that treatment with anti-CTLA-4 antibody between cisplatin cycles reduced tumour growth and increased survival [30]. The survival and reduced tumour growth was related to improved antitumor immune system responses, including an increase of the total number of T-cells, CD4+/CD8+ T-cells infiltrating into the tumour, and upregulation of IL-2, promoting T-cell activation. These positive preclinical outcomes lead to
clinical studies, highlighting CTLA-4 as the first immune checkpoint target for immunotherapy [31].

PD-1 is another immune checkpoint marker expressed on T-cell surfaces, which regulates inhibitory signals by its ligand PD-L1 and together with CLTA-4 represents the main promising cancer immunotherapeutic targets. Preclinical studies showed PD-L1 to be present on murine mesothelioma cells in vivo [32]. PD-L1 overexpression has been detected in response to interferon (IFN)-γ and T-cells in tumour-draining lymph nodes [32]. PD-L1 activation has been reported in mesothelioma specimens and is associated with worse prognosis [33-35]. From these results PD-L1 may represent a potential target for novel therapies in MM.

4. Interaction between the immune system and cancer

Cancer development is a multistep process in which different systems and molecular components are actively responsible for malignant properties [36]. Understanding the molecular mechanisms involved in the cancer microenvironment, in particular the role of stromal and inflammatory cells, is crucial to identify new targets as diagnostic biomarkers or to develop possible novel therapeutics. One particularly important component is the immune system [37]. This system is very tightly regulated by several cellular populations to activate both innate and adaptive immunity against pathogens. As they are an important component of adaptive immunity, T-cells require complex interactions of signals to be activated [38]. The main stimulatory regulation is mediated by signals between T-cell receptor (TCR) and peptide-major histocompatibility complexes (MHCs) on antigen-presenting cells (APCs). The interaction is amplified by an additional balance between co-stimulatory/co-inhibitory signals which play a central function in maintaining a sufficient immune response against pathogenic antigens and limited response against self-antigens, guaranteeing an immunologic homeostasis [38, 39].

Therefore, in normal conditions, the immune checkpoint receptors, such as CTLA-4, PD-1 and myeloid-derived suppressor cell (MDSC) interact with their ligands (B7, PD-L1) on various cells types, including APCs, regulatory T-cells (Tregs) resulting in reduced T-cell proliferation and functional activity. In the tumour microenvironment, immunologic tolerance has recently been shown to have major implications for the ability of tumours to survive and, in addition, immunosuppressive elements such as CTLA-4, and MDSC are implicated in this mechanism through a variety of pathways. It has been shown that in cancer, immune checkpoints (which exert a
negative effect on the immune system) are consistently activated, which leads to a loss of balance between co-stimulatory and co-inhibitory pathways, thus increasing tumour evasion [40].

5. **Immune response to MM**

Research into the role of the immune response in MM has been carried out, with the research highlighting the importance of the immune system in the development and promotion of MM. Once the asbestos fibres are inhaled, macrophages unsuccessfully attempt to metabolize and destroy this foreign body, leading to chronic inflammation [41]. Furthermore, reports indicate that survival may be increased by elevated numbers of CD8+ tumour-infiltrating lymphocytes [42]. Tregs, however, may infiltrate the tumour microenvironment and influence tumour growth. It has been shown that depletion of Tregs increases survival [43].

Features of immunogenicity in MM have been shown by studies that demonstrate that MM cells can act as Antigen Presenting Cells [44] and express Cancer Testis Antigens (CTA) noted as potential target for immunotherapy [45]. On the contrary MM cells interfere negatively with the immune activities dependent on IL-2 [46] and suppress the switching of T-cells towards a TH1 profile. Transforming growth factor-beta released by Purified Protein Derivative (PPD)-presenting malignant mesothelioma cells inhibits interferon-gamma synthesis by an anti-PPD CD4+ T-cell clone [47].

However, the first clinical studies related to these molecules have been disappointing [46, 48]. Therefore, the attention was focused on immunotherapeutic strategies using monoclonal antibodies (mAbs) that target receptors of immune cell surface or molecules expressed on tumour cells which regulate the anti-tumour activity of T-cells. Among these molecules, CTLA-4 and B7-H3 have been investigated in MM as immunotherapy [49] and have provided a rationale for their therapeutic use for this tumour.

6. **Clinical studies of tremelimumab in unresectable MM patients**

So far, a second-line therapy for MM is not established and there are no treatments that significantly prolong survival rate. The urgent need to find both high efficacy and well-tolerated therapy for refractory MM patients and the potential application of anti-CTLA-4 antibodies has led to the
design of an experimental study termed MESOT-TREM-2008 (NCT01649024) (Table 1) [50]. This study is an open-label, single-arm phase 2 trial where the clinical aspects and immunological activity of tremelimumab were evaluated in advanced malignant mesothelioma patients who failed the first-line treatment based on platinum-based regimen [50]. Twenty-nine highly selected patients (limited disease and very good performance status) were enrolled and received a dose of 15 mg/kg intravenously once every 90 days.

The primary endpoint was the proportion of treated patients who reached a complete or partial objective response according to the Response Evaluation Criteria in Solid Tumors (RECIST) for peritoneal malignant mesothelioma and modified RECIST criteria for MM. The secondary endpoint was to include the proportion of treated patients with disease control, progression-free survival (PFS), overall survival (OS), safety, and changes in lymphocyte phenotype induced by tremelimumab. Since only two patients (7%) achieved a durable partial response (one lasting 6 months and the other one lasting 18 months), the primary endpoint was not reached. Nine patients (31%) showed disease control, median PFS was 6.2 months and OS was 10.7 months. Serious side effects (3°-4° grade with one death due to gastrointestinal toxicity) were observed in four (14%) patients, with one demonstrating hepatic and pancreatic toxicity, and two developing gastrointestinal adverse events. The patient who achieved long-lasting partial response developed a Guillain-Barré-like syndrome. The persistent disease control and the long-lasting partial response together with an initial progressive disease are considered typical features of CTLA-4 treatment, as already reported in metastatic melanoma patients treated with ipilimumab [51]. Since these are non-typical patterns of clinical response, it is necessary to carefully evaluate these patterns before interrupting CTLA-4 because the clinical benefits may be delayed.

These findings lead to a redefinition of the response criteria for immune checkpoint therapies, defined immune-related response criteria (irRC) [52]. The main criteria, which distinct irRC from RECIST are requirement of confirmation of progression disease (PD) with two subsequent scans at least 4 weeks later, and new lesions are not necessary linked to PD but they need to be included in the total tumour burden (Table 2) [53]. Moreover, an increased number of circulating CD4+ICOS+ T lymphocytes, ICOS, expressed on the cell surface of activated T-cells, plays a role in T-cell expansion and survival, was associated with improved survival as reported in studies of ipilimumab or tremelimumab in melanoma and breast cancer [54, 55]. These findings suggest that circulating CD4+ICOS+ T lymphocytes may be a general tool to guide the use of anit-CTLA-4 mAbs.
Another phase 2 study for MM, called MESOTREM-2012 (NCT01655888) has explored a more intensive dosing schedule (Table 1). MESOTREM-2012 is an open-label, single arm, phase 2 trial for patients with unresectable, advanced MM (pleural or peritoneal) and progression disease after a platinum-based treatment regimen [56]. Tremelimumab was administrated for 1 hour intravenously at a dose of 10 mg/kg every 4 weeks for six doses, followed by maintenance dosing every 12 weeks. In this study, the primary endpoint was the proportion of treated patients who reached a complete or partial immune-related objective response according to the irRC criteria. The secondary endpoint was to include the proportion of treated patients with immune-related disease control, immune-related progression-free survival, OS, safety, and changes in lymphocyte phenotype induced by tremelimumab (see the comments on irRC in the “perspective” section for further details).

The primary endpoint was reached and four (14%) patients achieved an immune-related partial response, fifteen (52%) treated patients had a disease control (median of 10.9 months) and the median immune-related PFS was 6.2 months, whilst 11.3 months was the median overall survival. A modest increase of side effects (grade 1° and 2°) was observed compared to the previous study; two patients had grade 3 hepatic or gastrointestinal toxic effects. From this study, it was observed that the best population of T-cell in response to CTLA-4 blockade is ICOS-positive T-cell subset, which may represent a more accurate pharmacodynamics marker to predict better survival. These findings are currently under investigation in another randomised, double-blind, placebo-controlled, phase 2b trial study (NCT01843374) (Table 3).

Recently, it has been announced by AstraZeneca and MedImmune that tremelimumab as monotherapy does not improve survival and the primary endpoint is not reached (D4880C00003; NCT01843374) [57]. Further data regarding DETERMINE was presented at ASCO 2016, detailing that 571 patients were enrolled and that 81% had died throughout the duration of the study [58]. There was no statistically significant difference between treated patients and placebo patients, and side effects such as diarrhoea, loss of appetite, and development of rashes were seen at higher rates in treated patients than in placebo treatments [58]. This indicates a need for further research into the use of tremelimumab as a monotherapy as well as use in combination therapy to potentially achieve better clinical outcomes.

7. Combination Immunotherapy
It is hypothesised that combination immunotherapy could yield better clinical outcomes than monotherapy. Nivolumab and pembrolizumab, two PD-1 inhibitors, have been approved to treat patients with advanced and metastatic melanoma, as well as refractory non-small cell lung cancer [59]. Their use in mesothelioma is much less well-established, however there are some clinical trials that are currently underway or in recruiting. One trial (KEYNOTE-028) detailed preliminary findings on the use of pembrolizumab in MM and demonstrated that the drug was well-tolerated and had some anti-tumour effect [60]. However, for immunotherapy in general and in particular for immunotherapy for MM, much more research is needed, particularly due to failure of some studies to meet the primary endpoint [57]. The safety profile of combined immunotherapy is another aspect worth clinical assessment.

There have been several clinical studies combining anti-PD-1 and anti-CTLA-4 agents, such as one in advanced melanoma which identified that the combination therapy nivolumab and ipilimumab lead to an objective response of 53%, with up to 80% in tumour mass [61]. Another trial, using untreated patients with metastatic melanoma, demonstrated that nivolumab was more effective than ipilimumab (as was combination of nivolumab and ipilimumab against ipilimumab alone), and that for PD-1-negative patients combination therapy was more effective than monotherapy [62]. The combination of these targets has been approved for use in patients with BRAF wild-type metastatic or unresectable melanoma [59]. Again, the use of the combination therapy is less well-established in MM than for other cancer types, likely due to the slower development for the drugs as a monotherapy. However, anti-CTLA4 and anti-PD-L1 trials have been reported to be underway [63], and a study design for tremelimumab and durvalumab combination for MM was presented at ASCO 2016 (NIBIT-MESO-1) [64]. This study is in early stages and according to the ClinicalTrials.gov page it is still recruiting.

Ultimately immunotherapy, whether as a monotherapy or in combination, is still in its very early stages for mesothelioma. The recent setback of tremelimumab trial not meeting its study criteria [57] and the early stage of clinical trials signify the amount of work still needed in this area.

8. Key issues and future perspectives

Even though the potential of a future use of tremelimumab for MM looks has been largely jeopardized by the latest clinical outcome (which included some cases of very severe toxicity and a
lack of effect on survival), some more general comments should be considered before proceeding with immune checkpoint inhibitors for MM.

-Biological comments: recent studies on stroma and the microenvironment provide significant evidence which may suggest a potential therapeutic use of check point inhibitors for MM. Among other factors, T-cell exclusion looks to be likely to confer resistance of MM to this type of therapy [65].

-Clinical comments: so far only MM patients who are very fit and with limited disease have been recruited into checkpoint inhibitors trials; irRC as a response criteria are very arguable and unlikely to be reliable for this tumour whose thoracic extension and actual shrinkage after treatment are very hard to measure. Patient stratification is a very relevant issue that should be properly addressed but is still at early stages; as the study population is getting larger, our knowledge on acute and chronic toxicity is unravelling more side effects that should be now considered. Lastly, though probably the most important aspect, is the real impact on survival of these treatment. Other than some anecdotal very good responses, no positive impact on survival has been achieved with these treatments so far.

-Cost/benefit: all the above comments cannot be considered without bearing in mind the financial cost that the use of these drugs implies and that, at the moment, such cost cannot be justified by the clinical results.

-Future indication: when the toxicity results reach acceptable levels, the effect on survival of combined checkpoint inhibitors and of checkpoint inhibitors along with chemotherapy are worth investigating. Addressing checkpoint inhibition and the microenvironment is clearly a further fascinating, although still speculative, option. Eventually an adjuvant role of these treatments could be also cautiously hypothesized. All of the above biological and clinical comments are clearly in place also for these potential approaches.

Financial disclosure
Nonfinancial— No relevant nonfinancial relationship exists.
Executive summary

- It has been shown that in cancer, immune checkpoints (which exert a negative effect on the immune system) are consistently activated, which leads to a loss of balance between co-stimulatory and co-inhibitory pathways, thus increasing tumour evasion.

- The role of immune responses has been deeply investigated in MMe, highlighting the importance of the immune system in developing and promoting MMe.

- CTLA-4 is an immune checkpoint receptor which regulates inhibitory signals to guarantee an immunological homeostasis.

- Blockage of the CTLA-4 receptor allows T-cell activation and tumour-specific T-cells to exert cytotoxic effects on tumour cells.

- Tremelimumab is a fully human monoclonal antibody used in clinical trials for unresectable MMe patients.

- Conclusion: biological and clinical considerations may rule out the use of Tremelimumab as single agent for MMe and the use of immune checkpoint inhibitors for MMe is still largely questionable and not supported by current evidence.

Figure 1. CTLA-4-B7 axis is a co-inhibitor signal for T-cell activation. Effector T-cell activation is mediated by a main interaction between TCR and MHC which is amplified by a co-stimulatory signal (B7/CD-28 binding) (A). Self-tolerance and immunologic homeostasis is regulated by a negative signal between CTLA-4 with B7 in normal cells (B). Anti-CTLA-4 antibodies inhibit CTLA-4 activity (C).
Table 1. Clinical trials with tremelimumab in advanced malignant mesothelioma patients.

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<td>NCT01649024</td>
<td>The immunologic and clinical activity of Tremelimumab as second line treatment</td>
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<tr>
<td>MESOT-TREM-2012</td>
<td>NCT01655888</td>
<td>The efficacy of a more intensive schedule of treatment with Tremelimumab as second line treatment</td>
<td>29</td>
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<tr>
<td>D4880C00003</td>
<td>NCT01843374</td>
<td>Randomized, double-blind, parallel-group study (Tremelimumab versus placebo as second/third line)</td>
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Table 2. Overall response of immune related response criteria (irRC) Information taken from Wolchok and colleagues [52].

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<tr>
<td>Complete Response (CR)</td>
<td>Disappearance of all lesions in two consecutive observations not less than four weeks apart</td>
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<tr>
<td>Partial response (PR)</td>
<td>A ≥50% decrease in tumour burden compared with baseline in two observations at least four weeks apart</td>
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<td>Stable Disease (SD)</td>
<td>A 50% decrease in tumour burden compared with baseline cannot be established nor 25% increase compared with nadir</td>
</tr>
<tr>
<td>Progression Disease (PD)</td>
<td>At least 25% increase in tumour burden compared with nadir (at any single time point) in two consecutive observations at least four weeks apart</td>
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44. Mutti L, Valle MT, Balbi B et al. Primary human mesothelioma cells express class II MHC, ICAM-1 and B7-2 and can present recall antigens to autologous blood lymphocytes. *Int J Cancer* 78(6), 740-749 (1998).


