The Role of B and T Lymphocyte Attenuator in Respiratory System Diseases

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B and T lymphocyte attenuator (BTLA), an immunomodulatory molecule widely expressed on the surface of immune cells, can influence various signaling pathways and negatively regulate the activation and proliferation of immune cells by binding to its ligand herpes virus entry mediator (HVEM). BTLA plays an important role in immunoregulation and is involved in the pathogenesis of various respiratory diseases, including airway inflammation, asthma, infection, pneumonia, acute respiratory distress syndrome and lung cancer. In recent years, some studies have found that BTLA also has played a positive regulatory effect on immunity system in the occurrence and development of respiratory diseases. Since severe pulmonary infection is a risk factor for sepsis, this review also summarized the new findings on the role of BTLA in sepsis.

Keywords: BTLA, airway inflammation, lung cancer, asthma, sepsis

INTRODUCTION

Nowadays, more and more studies focus on the immunological pathogenesis of diseases. Immune dysfunction plays a pivotal role in the development of inflammation, infection and tumor. Many immunomodulatory chemicals and targeted drugs have been used in clinical practice. As a co-signaling molecule, B and T lymphocyte attenuator (BTLA) plays an important role in immunoregulation and is involved in the pathogenesis of various respiratory diseases. In this review, we discuss the biological characteristics of BTLA and explore their role in respiratory diseases.

THE BIOLOGICAL FUNCTION OF BTLA

BTLA, a co-inhibitory molecule similar to cytotoxic T lymphocyte antigen-4 (CTLA-4) and programmed death 1 (PD-1), belongs to the immunoglobulin superfamily, and mainly negatively regulates the activation and proliferation of immune cells (1, 2). BTLA is widely expressed on the surface of various immune cells, such as B cells, T cells, monocytes, macrophages, dendritic cells (DCs), natural killer cells (2, 3). At all phases of T cell differentiation including naive T cell, BTLA expression exists (4, 5). BTLA is a transmembrane glycoprotein. Its extracellular domain is immunoglobulin domain, and intracellular domain contains three conserved motifs: one is the proximal motif which have sequence YDND, an immunoreceptor tyrosine-based inhibitory motif
BTLA in Respiratory System Diseases

BTLA IN AIRWAY INFLAMMATION AND ASTHMA

Allergic airway inflammation can cause immune response in the lung, which is mediated by Th2 cells and their cytokines such as IL-4, IL-5, and IL-13, and subsequently cause epithelial damage and airway hyperreactivity (19). As an immunoregulatory molecule, BTLA is involved in the development of allergic airway inflammation (Table 1). Deppong C et al. found that after allergens inhalation, the airway inflammation of the sensitized wild type mice reached to its peak by day 3, and resolved by day 10, while the airway inflammation of the mice deficient in expressing BTLA could last as long as 15 days, indicating that BTLA can shorten the duration of allergic airway inflammation and give proper termination to the acute inflammatory response (20). Another research focused on the role of BTLA in the proliferation, recruitment, and survival of T cells in response to inhaled allergens in BTLA-deficient mice and wild type mice. Decreased cell death of T cells was found in BTLA deficient mice, whereas proliferation and recruitment of T cells to the lungs remained unaffected, indicating BTLA signaling is a key determinant of the inflammatory response in the lung (21). Apart from that, BTLA is involved in suppression of the induction of allergic airway inflammation. Tamachi et al. found IL-5 production and eosinophilic inflammation were increased in the airways of BTLA deficient mice after the antigen inhalation (22). The above research results showed that BTLA could regulate the death of T cells in lung and inhibit the aggregation of eosinophils induced by endobronchial antigens. However, the precise mechanism is still unclear and no information on the role of BTLA in allergic diseases in humans is available. Further analysis is needed to reveal the role of BTLA in the biologic basis of eosinophil-mediated allergic diseases. Considering that asthma is associated with intrabronchial aggregation of eosinophils and disproportion of T cell subsets, BTLA may be correlated with the pathogenesis of asthma. A research with children found that the variation of multi-loci on BTLA gene could influence serum IgE levels (23), which indirectly indicated that BTLA might be associated with asthma. These studies indicated that BTLA may serve as a novel target for the therapeutic intervention.

BTLA AND INFECTION

Antigen-specific T cells are crucial for the anti-infective effect. Recent studies have shown that BTLA also plays a role in immune responses against infectious pathogens. Since BTLA is widely expressed on the surface of immune cells, its relation to immune response of infection including innate immune and adaptive immune has been a hot topic. A research indicated that BTLA was critical for negatively regulating early host immunity against intracellular bacteria. Compared with wild type mice, HVEM and BTLA deficient mice were more resistant to listeriosis. Blocking BTLA signaling pathway could promote early removal of bacteria. Stimulated by the Listeria, innate immune cells of BTLA deficient mice secreted significantly more proinflammatory factors, which indicated that BTLA played an important regulatory role in early host innate immune response against infection (24). BTLA also negatively regulates immune response to virus. Cytomegalovirus (CMV) infection could induce high expression of BTLA on virus specific CD8⁺ T cells. Using antibody to block BTLA in vitro could facilitate the proliferation of virus specific CD8⁺ T cells (25). On the contrary, some studies found that the expression of BTLA helped fight against infection.
Marcos W. Steinberg et al. found that in BTLA and HVEM deficient mice and mice with an BTLA-HVEM blockade, the number of antigen specific CD8+ T cells was reduced after bacterial infection. This result suggests that BTLA-HVEM signaling pathway does not restrict to inhibitory signaling transmission, BTLA can promote the survival of antigen specific CD8+ T cell to fight against bacterial infection through HVEM dependent signal pathway (26). Similar results were also found in viral infection. The numbers of the effector CD8+ T cells and memory CD8+ T cells were reduced after BTLA or HVEM deficient mice infected with vaccinia virus. HVEM-BTLA signaling could promote the differentiation of memory CD8+ T cells to defend viral infection (27). The positive effect of the signaling pathway may be related to tran-interaction of BTLA with HVEM, while cis-interaction shows negative effect. Coronavirus disease 2019 (COVID-19), which is caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), is globally pandemic. There were some researches showing the relation between BTLA and COVID-19. Both Christoph Schultheiß et al. and Narjes Saheb Sharif-Askari et al. found that BTLA was upregulated compared to that in controls (28, 29). Marissa Herrmann et al. found that the level of BTLA on CD8+ T cell decreased in COVID-19, but not as strong as in healthy controls, and the expression of BTLA on transitional memory and effector memory CD8+ T cells in COVID-19 was higher compared to healthy controls (30).

The researches on the role of BTLA in pulmonary infection mainly focused on tuberculosis (TB) infection (Table 2). The chronic infection of Mycobacterium tuberculosis (Mtb) indicates the protective (escaping) strategies to avoid clearance by the innate and adaptive immune responses (36). Shen et al. found that BTLA was upregulated on circulating CD4+ and CD8+ T cells of pulmonary TB patients. The level of BTLA expression was dynamically changed with the increase of TB bacillary load, suggesting that BTLA could be used as a useful marker reflecting immune function as well as disease progression (33). Wang et al. analyzed the role of BTLA in antigen presenting cells (APCs) and found that BTLA was highly expressed in CD11c-expressing APCs in patients with active pulmonary tuberculosis (ATB). The BTLA-expressing CD11c APCs showed decreased capacity to stimulate allogeneic T cell proliferation which was associated with low

### TABLE 1 | The function of BTLA in acute allergic airway inflammation.

| Study. Year | Specimen | Immune cells | Subjects | Main findings | Reference |
|-------------|----------|--------------|----------|--------------|-----------|
| Deppong et al., 2006 | BALF | BTLA+ WT mice | 68 ATB patients 40 healthy controls | 1. BTLA expression on CD8 T cells is decreased in ATB patients. 2. BTLA expression on CD4 T cells is decreased in ATB patients. | (31) |
| Tamachi et al., 2007 | Spleen | BTLA+ WT mice | 52 ATB patients 15 healthy controls | 1. The frequencies of BTLA positive CD11c APCs in ATB patients were higher than that in healthy controls. 2. BTLA-expressing CD11c APCs in ATB patients show low capacity to stimulate T cell proliferation. | (32) |
| Deppong et al., 2008 | BALF | BTLA+ WT mice | 86 ATB patients 40 healthy controls | The levels of BTLA expression were upregulated on peripheral CD4+ and CD8+ T cells of ATB patients and associated with disease progression. | (33) |
| Cai et al., 2019 | Pleural effusions | mDCs | 20 tuberculosis pleurisy patients 15 healthy controls | 1. Co-expression of BTLA and B7-H4 on myeloid dendritic cells DCs (mDCs) in peripheral blood and pleural effusions of pleural TB patients was significantly higher than in the control group. 2. High expression of BTLA and B7-H4 promoted a high level of CD83 and HLA-DR, which had a negative regulatory effect on mDCs on anti-TB immunity. | (34) |
| Zhang et al., 2020 | Peripheral blood samples | mDCs and pDCs | 73 ATB patients 35 healthy controls | 1. ATB patients exhibited higher expression of BTLA in mDCs and pDCs subsets than healthy controls. 2. TB-driven BTLA expression in DCs impairs the expression of functional DC surrogate markers and suppress the ability of DCs to induce anti-TB Th17 and Th22 response while promoting Th2 and Foxp3+ Tregs. | (35) |

**APCs, antigen presenting cells; ATB, active pulmonary tuberculosis; BTLA, B and T lymphocyte attenuator; mDCs, myeloid dendritic cells; Mtb, mycobacterium tuberculosis; pDCs, plasmacytoid DCs; TB, tuberculosis.**
expression of HLA-DR and less IL-6 secretion in ATB patients (32). An extension study showed that TB-driven BTLA expression in DCs could affect their biological characteristics and immune functions, which was associated with an increased capacity to produce IL-4 and TGF-β and a decreased capacity of DCs to produce the key cytokine IL-12, and to induce T cell proliferation and differentiation into Th subsets, resulting in altered anti-TB immune responses and immunity (35). An analogous finding showed that high co-expression of BTLA and B7-H4 on myeloid dendritic cells (mDCs) in peripheral blood and pleural effusions of pleural TB patients promoted a high level of CD83 and HLA-DR, which had a negative regulatory effect on mDCs and anti-TB immunity (34). In contrast to the up-regulation of BTLA expression in circulating CD4+ and CD8+ T cells, APCs, and DCs, the expression of BTLA was decreased in αβ T cells of active pulmonary tuberculosis patients and anti-tuberculosis drugs induced BTLA expression along with bacterial clearance. BTLA expression on αβ T cells was associated with protective immune memory in ATB patients against Mtb infection (31). Unlike the role of BTLA in negative regulation of immune responses, this result indicates that BTLA is involved in pathogen clearance.

**BTLA AND SEPSIS**

Inflammatory responses play a critical role in the pathogenesis of pneumonia, and the intensity of these responses often determines the severity of the disease. Severe pulmonary infection is a risk factor for sepsis. BTLA signaling can induce several immune responses such as immune tolerance, immunosuppression, and immune escape. Previous researches have demonstrated that BTLA plays a role in regulating the immune response in sepsis (Table 3). Nicholas J. Shubin et al. found that the number of BTLA and HVEM expressing macrophages, dendritic cells, neutrophils increased in the original infection site of septic mice. BTLA deficient septic mice showed higher survival rate than wild type septic mice (37). Another research showed similar results that BTLA could suppress LPS induced endotoxic shock by suppressing cytokine production from LPS-stimulated dendritic cells and macrophages (14). However, Cheng et al. found different phenomena. Treating septic mice with anti-BTLA antibody, cytokines and inflammatory cells increased in the original site of infection, and the mice exhibited more severe organ impairment and lower survival rate (40). Differences also exist among clinical researches. A research found that the level of soluble BTLA in the serum of septic patients was much higher than that of ICU non-septic control and healthy control, and the level was associated with Sequential Organ Failure Assessment (SOFA) score, which is calculated by various indicators such as the oxygenation index, the Glasgow coma scale, the level of platelet, creatinine and so on. The level of sBTLA in 28 days sepsis non-survivors was significantly higher than in survivors (41). Similar result was

**Table 3** | The function of BTLA in sepsis.

| Study | Year | Specimen | Subjects | Main findings | Reference |
|-------|------|----------|----------|---------------|-----------|
| Shubin et al. 2012 | Peripheral blood | BTLA+/CD4+ mice | WT mice | 1. The number of infiltrating BTLA- and HVEM-expressing macrophages, inflammatory monocytes, mature and immature DCs, and neutrophils increased in the peritoneum in mice with acute experimental sepsis induction. 2. BTLA and HVEM monocytes in peripheral blood and HVEM granulocytes were increased in septic ICU patients. 3. BTLA can serve as makers to predict the occurrence of sepsis. | (37) |
| Shubin et al. 2013 | Peripheral blood | BTLA+/CD4+ mice | WT mice | 1. The septic ICU patients had a higher percentage of BTLA+ CD4+ lymphocytes in peripheral blood compared with critically ill non-septic individuals. 2. BTLA expression in circulating CD4+ T-cell and B-cell increased in septic mice. 3. II patients with CD4+ T-cells expressing greater than 80% BTLA+ had longer hospital stays. | (38) |
| Shao et al. 2015 | Peripheral blood | 286 sepsis patients | 50 healthy controls | 1. BTLA+/CD4+ T cells was high expressed in healthy volunteers and was reduced in severe sepsis and septic shock patients. 2. The percentage of BTLA+/CD4+ T cells was lower in non-survivors than that in survivors. | (39) |
| Cheng et al. 2016 | Peripheral blood | Sepsis mice model | 101 patients with sepsis | 1. BTLA expression is elevated on innate immune cells in mice model of hemorrhagic shock/sepsis. 2. Anti BTLA antibody treatment increased cytokine/chemokine levels and inflammatory cells recruitment, aggravated organ injury and elevated these animals’ mortality. | (40) |
| Lange et al. 2016 | Peripheral blood | Soluble BTLA levels in plasma were higher in the sepsis cohort and is associated with severity of disease. | 28 ICU controls | (41) |
| Arens et al. 2016 | Peripheral blood | BTLA was upregulated in CD4+ cells of sepsis survivors. | 31 healthy controls | (42) |
| Kobayashi et al. 2018 | Spleen | BTLA+/CD4+ mice | WT mice | 1. BTLA+ mice are more susceptible to LPS-induced endotoxic shock. 2. BTLA inhibit LPS-induced cytokine production in dendritic cells and macrophages. 3. Anti-BTLA antibody save mice from LPS-induced endotoxic shock. | (14) |

**BTLA, B and T lymphocyte attenuator; BTLA−/−, BTLA-deficient; WT, wild-type; LPS, lipopolysaccharide.**
found by Sean F. Monaghan et al., and sBTLA can predict the
diagnosis of sepsis (43). Nicholas J Shubin et al. exhibited that
in the peripheral blood from ICU patients with sepsis, the
proportion of BTLA expressing CD4+ T cells increased. In
critically ill patients without sepsis, if over 80% of the CD4+ T
cells expressed BTLA, they developed nosocomial infections
more easily and had longer hospital stays (38). The BTLA
density on the surface of peripheral blood CD4+ T cells was
upregulated in sepsis survivors compared to healthy controls
(42). Differently, Rui Shao et al. found that in severe sepsis and
septic shock patients, the proportion of peripheral blood
BTLA+/CD4+ T cells was significantly reduced compared with
healthy volunteers, and that ratio was lower in septic non-
survivors compared to septic survivors (39). The differences
among above researches may be due to the timing of entry
points. The expression of BTLA at different stage of sepsis may
have different clinical effects. At the early stage of sepsis, which
is the proinflammatory stage, the expression of BTLA may
increase along with the enhanced inflammatory reaction, so as
to protect organs from inflammatory storm. While at the anti-
inflammatory stage, high expression of BTLA inhibits the
activation of immune cells, and excessive immune
suppression may lead to a secondary infection and bad prognosis.

**BTLA in Pneumonia and Acute Respiratory Distress Syndrome**

Inflammatory responses are involved in the
immunopathogenesis of pneumonia, especially severe
pneumonia disease. The intensity of these inflammatory
responses often determines the severity of the disease (44).
One study focused on pneumonia demonstrated that BTLA
protein expression was mainly present in the bronchial
epithelium and inflammatory cells in patients with severe
community-acquired pneumonia (CAP), suggesting that BTLA
might be involved in host protection. The percentages of
circulating BTLA+CD4+ lymphocytes were significantly higher
in patients with severe CAP and in mice with lipopolysaccharide
(LPS)-induced acute lung inflammation than in control groups.

Increasing BTLA expression via either the administration of
dexamethasone or the agonistic anti-BTLA antibody 6A6
attenuates LPS-induced acute lung inflammation in mice
(Table 4) (45). BTLA may be involved in regulating the
immune response in patients with severe CAP, affecting the
outcome of this disease.

Severe pneumonia may cause acute respiratory distress
syndrome (ARDS). ARDS is an organ failure syndrome caused
by inappropriate and uncontrolled inflammatory response. The
main pathophysiological feature of ARDS is inflammatory
cytokine storm caused by overactivation of the immune system
(47, 48). The expression of BTLA was increased on the surface of
alveolar macrophages (AMs) and pulmonary CD4+ lymphocytes
of ARDS rats (46). The treatment with lung-resident
mesenchymal stem cells (LRMSCs), a promising candidate for
ARDS therapy by regulating excessive inflammatory responses,
can increase the expression of BTLA on immune cells. When
BTLA expression was knocked down by siRNA, the
immunoregulatory effects of LRMSCs were partially abolished,
indicating that BTLA is involved in the immunoregulatory
process operated by LRMSCs (Table 4) (46). Thus, BTLA may
serve as a target for ARDS treatment.

**BTLA and Lung Cancer**

The evasion of immune attack and formation of an immune
suppressive environment within tumor exist during the entire
process of cancer development (49). Immune checkpoint
molecules can create immunosuppressive conditions in various
cancers to impact tumorigenesis (50). Previous researches have
shown the up-regulation of BTLA in gastric cancer (51),
pancreatic cancer (52) and lymphocytic leukemia (53) and
BTLA overexpression has been found to be associated with an
immunosuppressive microenvironment (54). The blockade of
the immunoinhibitory HVEM-BTLA/CD160 pathways may
result in sustained tumor regression (55). One study found
that the expression of BTLA on intratumoral CD8+ T cells was
enhanced along with the progression of disease, and the co-
expression of BTLA and other co-inhibitory molecules could

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**Table 4** | The function of BTLA in severe CAP and ARDS.

| Study. Year | Specimen | Subjects | Main findings | Reference |
|------------|----------|----------|--------------|----------|
| Zhou et al., 2016 | Peripheral blood BALF (only in mice) Mucosal biopsy specimens | 11 patients with severe CAP 10 healthy controls Mice with LPS-induced acute lung inflammation Control mice | 1. The percentages of circulating BTLA+CD4+ lymphocytes were significantly higher in patients with severe CAP and in mice with LPS-induced acute lung inflammation than in the control groups. 2. BTLA was mainly expressed in the bronchial epithelium and inflammatory cells. | (45) |
| Cheng et al., 2020 | Lung | An ARDS group of rats A PBS control group of rats An ARDS + LRMSCs group of rats | 1. The expression of BTLA was increased on the surface of alveolar macrophages (AMs) and pulmonary CD4+ lymphocytes of ARDS rats. 2. BTLA is involved in the immunoregulatory process operated by LRMSCs | (46) |

ARDS, acute respiratory distress syndrome; BALF, bronchoalveolar lavage fluid; BTLA, B and T lymphocyte attenuator; CAP, community-acquired pneumonia; LPS, lipopolysaccharide; LRMSCs, lung-resident mesenchymal stem cells.
inhibit T cell function (56). Similarly, the expression of BTLA was increased on CD4\(^+\) T cells and CD8\(^+\) T cells isolated from pleural effusion of lung cancer patients, indicating BTLA might mediate a negative cosignal for local immune response (57). Furthermore, BTLA was also expressed in tumor cells of non-small cell lung cancer (NSCLC) patients and the BTLA levels were significantly higher in patients with lymphatic metastasis and high tumor pathological stage (58). Those findings indicate that lung cancer can affect the body’s immune status through BTLA. A research in mice that received subcutaneous implantation of lung cancer cells showed that the expression of BTLA on CD4\(^+\) T cells and CD8\(^+\) T cells increased and the number of these T cells increased as well, while BTLA\(^+\)/CD8\(^+\) T cells produced less IL-2 and TNF. The results indicated that tumor could induce enhanced expression of BTLA by T cells, impair T cell functions, thus led to systemic immunosuppression state (59). This may be the reason why tumor can escape from immune surveillance, and make tumor-bearing individuals more susceptible to infections (60). Moreover, a research showed that in lung adenocarcinoma that displayed an epithelial-mesenchymal transition (EMT) phenotype, BTLA expression was elevated in mesenchymal tissues, indicating that BTLA might influence the EMT of tumor by changing the inflammatory tumor microenvironment, then influence tumor metastasis and drug resistance (61). As for the impact on prognosis, Li et al. found high BTLA expression might predict the progression and poor prognosis of NSCLC. Patients with positive BTLA expression had a shorter relapse-free survival (RFS) than those with negative BTLA expression (58). A pharmacogenetic study suggested that a BTLA polymorphism with potential function to modify miRNA binding sites (rs76844316) was connected to the occurrence and prognosis of lung cancer (62). BTLA may be a novel therapeutic target for cancer immunotherapy (Table 5).

**DISCUSSION**

BTLA plays an important role in immunoregulation and is involved in the pathogenesis of various respiratory diseases (Figure 1). In spite of its importance in regulating immunity, the HVEM-BTLA signaling in respiratory system diseases has not been sufficiently analyzed. One reason is that BTLA does not merely serve as an immune suppression role in respiratory system diseases. In many circumstances, BTLA can promote immunity and fight against infection. BTLA contains structure of promotive function (63), so it may produce immune enhancement signals during signal transduction. Besides, the binding of BTLA to its ligand HVEM can form a bi-directional signal system. BTLA and HVEM can act as ligand and receptor for each other, delivering different signals. When BTLA binds to HVEM as a ligand, it generates positive immune regulation (9, 64). For example, the BTLA involvement can induce HVEM-mediated NF-xB activation, which is important for the induction of pro-inflammatory and cell survival genes (65). In addition, BTLA lays in a complicated network of immune modulation and signal transmission. Researchers separating one pathway from the network may not be comprehensive enough. Since HVEM and BTLA are widely expressed by many cell types, the exact regulatory mechanism in different immune contexts need to be carefully determined. So far, most researches merely find how the level of BTLA changes in different respiratory diseases. The underlying mechanism is still unknown. How HVEM-BTLA signaling regulates immunity and influences the pathogenesis of respiratory diseases need to be elucidated. More researches on mechanisms should be conducted. Based on current research, the level of BTLA may be used as an indicator of disease severity, and may predict the prognosis. Anti-BTLA antibody has been used in animal experiments to treat severe community-acquired pneumonia and epithelial ovarian carcinoma (45, 66). Antibodies in researches exert different effects, either agonistic or antagonistic.

| Study, Year | Specimen | Subjects | Main findings | Reference |
|-------------|----------|----------|---------------|-----------|
| Wang et al., 2006 | Peripheral blood and pleural fluid | 6 patients with lung cancer 6 healthy controls | The expression of BTLA was increased in the CD4\(^+\) and CD8\(^+\) T cells of pleural fluid of patients with lung cancer. | (57) |
| Thommen et al., 2015 | Fresh tumor tissues and malignant effusions | 32 patients with NSCLC | BTLA was generally expressed at a low percentage of tumor-infiltrating CD8\(^+\) T cells. | (56) |
| Mittal et al., 2015 | Spleen | Mice that received subcutaneous implantation of lung cancer cells wild-type mice 439 patients with lung adenocarcinomas from three clinical databases | 1. The frequencies of BTLA\(^+\) cells in both the CD4\(^+\) and CD8\(^+\) T cell compartments were increased in mice with localized cancer relative to non-cancer controls. 2. BTLA\(^+\)/CD8\(^+\) T cells in cancer mice exhibited reduced IL-2 and TNF. BTLA were elevated in mesenchymal lung adenocarcinoma. | (59) (61) |
| Lou et al., 2016 | Paraffin-embedded tissues | 87 patients with stage I–III NSCLC | | |
| Li et al., 2020 | Paraffin-embedded tissues | 169 patients with lung cancer 300 healthy controls | 1. BTLA was expressed in tumor cells in 35 patients with NSCLC (40.2%). 2. BTLA levels were significantly higher in NSCLC patients with lymphatic metastasis and high tumor pathological stage. 3. Patients with positive BTLA expression had a shorter relapse-free survival (RFS) than those with negative BTLA expression. BTLA rs1982809 AG genotype carriers had a higher risk of developing lung cancer when compared to AA genotype carriers in Tunisian population. | (58) (62) |

BTLA, B and T lymphocyte attenuator; NSCLC, non-small cell lung cancer.

TABLE 5 | The function of BTLA in lung cancer.
which may due to HEVM-BTLA bi-directional signal system. An anti-BTLA monoclonal antibody has been approved for clinical trial by FDA (68). Treatment of respiratory diseases by anti-BTLA antibody must be on the road, more and more application researches will be conducted.

**AUTHOR CONTRIBUTIONS**

ZD and YZ reviewed the literature and drafted manuscript. PC edited and revised manuscript. ZZ was responsible for the conception. All authors contributed to the article and approved the submitted version.

**FUNDING**

This work was supported by Research Project of Health Commission of Hunan Province (202103021536, 20200136, 20200121), National Natural Science Foundation of China (81903111), as well as Natural Science Foundation of Hunan Province (2020JJ8077).

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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