Opportunities and Challenges for Gut Microbiota in Acute Leukemia

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Acute leukemia (AL) is a highly heterogeneous hematologic malignancy, and although great progress has been made in the treatment of AL with allogeneic hematopoietic stem cell transplantation (Allo-HSCT) and new targeted drugs, problems such as infection and GVHD in AL treatment are still serious. How to reduce the incidence of AL, improve its prognosis and reduce the side effects of treatment is a crucial issue. The gut microbiota plays an important role in regulating disease progression, pathogen colonization, and immune responses. This article reviews recent advances in the gut microbiota and AL pathogenesis, infection, treatment and its role in allo-HSCT.

Keywords: gut microbiota, acute leukemia, pathogenesis, GVHD, drugs, immunity, allo-HSCT

INTRODUCTION

Acute leukemia (AL), a kind of hematologic malignancy, mainly includes acute myeloid leukemia (AML) and acute lymphoblastic leukemia (ALL) (1). In 2018, a global survey of 36 cancers in 185 countries indicated that leukemia ranks fifteenth in cancer incidence and tenth in cancer deaths (2). The most common cancer in children is AL (3). More than 80% of children with ALL are long term survivors, compared with 20% to 40% of adults (4). In AML, the five-year survival rate is 40% for patients younger than 60, and only 10-20% for patients older than 60 (5). Cytotoxic chemotherapy, ionizing radiation, genetic mutations, benzene, and some blood system diseases (such as myelodysplastic syndrome, paroxysmal nocturnal hemoglobinuria) can lead to AL (6, 7). The specific pathogenesis of AL is not fully understood, and some types of AL are dangerous, ineffective and have a high mortality rate. Immune system plays an important role in the pathogenesis, progression and treatment of AL. AML patients have higher T regulatory cells (Tregs) and CD8+ T cells compared to normal donors (8). However, AML patients have T-cell dysfunction, which is associated with immune escape of the disease (9). Moreover, Tregs are a major contributor to defective immune responses (10). In AML patients, increased Tregs can suppress effector T cells, and the more Tregs, the worse the treatment effect (11). Treatment with chimeric antigen receptor T-cell therapy is highly effective in patients with ALL, which is also associated with the immune system (12).

There are four major colonization sites of microorganisms in the human body: oral, vagina, skin, and gut (13). Gut microbiota colonize the newborn following birth (14). The gut microbiota is composed of native flora and transient flora in food intake (15). It is estimated that each individual’s gut microbiome includes more than 100 different species (15). Gut microbiota has recently been recognized as an important factor in regulating disease progression, pathogen colonization, and immune responses (1, 13, 16), and the pathogenesis, progression, and treatment of AL have been
linked to the immune system. Gut microbiota has also been linked to diabetes, Alzheimer’s disease, tumor and so on (17–19). This review summarizes the progress of the interaction between gut microbiota and AL.

THE ROLE OF GUT MICROBIOTA IN THE PATHOGENESIS OF AL

The pathogenesis of AL is unclear. Greaves M propose a model for ALL, in which some children are born with pre-leukemic clones, but acquired infections cause key genetic mutations that develop into ALL (20). Some studies have found that newborns carry leukemia-related mutations, such as TCF3-PBX1 and ETV6-RUNX1 (21, 22). Exposure to infection promotes the occurrence of ALL in newborns carrying these genes (23, 24). In mice with PAX5 or ETV6-RUNX1, ALL was caused by a disruption in the microbiome caused by antibiotic treatment early in life, and the gut microbiome of PAX5-carrying mice differed from the wild type (25). A study enrolled 81 subjects, including 58 ALL patients and 23 healthy controls, using 16S rRNA quantitative arrays and bioinformatics analysis to examine their stool specimens (26). Many species of bacteria appeared significantly different in stool specimens from ALL patients compared to healthy controls, such as a significant enrichment of Bacteroides claus in ALL (26). Whether the gut microbiota disturbance can cause AL in humans needs further study.

Alterations in some gut microbiota species and metabolites may promote the development of leukemia. In a mouse model of leukemia, the proportion of bacteria with the function of converting dietary flavonoids in the small intestine of leukemic mice was increased (27). It has been suggested that maternal intake of bioflavonoids may lead to infant and young child leukemia (28). A national case-control study conducted in Denmark and found that infections of the gastrointestinal tract were associated with an increased risk of AML (29). So why does the occurrence of a gastrointestinal infection lead to an increased risk of developing AML? We assume that the imbalance of gut microbiota due to gastrointestinal infections causes immune disorders that lead to AML, but more studies are still needed to verify this.

The risk of AL increased significantly during childhood or adolescence after the cumulative dose of ionizing radiation (30). Ionizing radiation is the accepted cause of leukemia. A study found that mice that survived high doses of radiation had unique gut microbiota, with Lachnospiraceae and Enterococcaceae as the most abundant bacteria (31). Then, they collected stool samples from 21 leukemia patients who had received total body radiation before hematopoietic stem cell transplantation, patients with shorter duration of diarrhea (diarrhea less than 10 days) had more Lachnospiraceae and Enterococcaceae in their feces than patients with longer duration of diarrhea (diarrhea more than 10 days) (31). Lactobacillaceae also increased in patients with shorter duration of diarrhea, but there was no statistical significance (31). Bone marrow and small intestinal epithelial cells are highly sensitive to ionizing radiation, and they are the main site of radiation injury (32). Short-chain fatty acids (SCFAs), a metabolite of gut microbiota, reduces DNA damage and reactive oxygen species release in blood and gastrointestinal tissues during radiation damage (31). Fecal microbiota transplantation (FMT) improved the survival rate of irradiated animals, increased white blood cells, and improved gastrointestinal function (33). The gut microbiota plays a key role in host defenses against radiation, as well as protecting the hematopoietic and gastrointestinal systems.

Benzene is a widespread air pollutant and a component of gasoline, industrial emissions, automobile exhaust and tobacco smoke (34). Traffic-related benzene exposure can cause acute myeloid leukemia in children (35). A study of benzene exposure in mice showed that benzene exposure altered the gut microbiota, with increased level of Actinobacteria, and Helicobacter were observed in the stools of mice exposed to high doses of benzene, and increased level of Actinobacteria were significantly and negatively correlated with basic blood cells (white blood cell, red blood cell, and hemoglobin levels) (36). Whether gut microbiota plays a role in benzene causing leukemia remains to be studied.

It is well known that naïve helper T cells (Th0) can be transformed into Th1, Th2, Treg and Th17 cells etc. Th1 produces the cytokine interferon gamma (IFN-γ) that exerts anti-tumorigenic effects in the tumor microenvironment, while Th2/Treg produces the cytokines interleukin 4 (IL-4), IL-5, IL-10, etc. that mediate pro-tumorigenic effects (17). In some solid tumor models, depletion of the gut microbiota leaded to a significant increase in IFN-γ-producing T cells and a decrease in IL-17A and IL-10-producing T cells, thereby reduced tumor burden. However, this phenomenon was not present in mice lacking mature T and B cells, suggesting that the action of the gut microbiota on tumor cells required the active involvement of adaptive immunity (17). In innate and adaptive immunity, microbiota plays an important role in its maturation, development and functions (37).

There is no doubt that the development of AL is related to the dysregulation of the human immune system, and the gut microbiota also regulates the human immune system, which has been shown to be closely related to the development of pancreatic cancer, colon cancer and other tumors (38, 39). Whether the gut microbiota is also involved in the development of AL by modulating human immunity needs to be studied in more depth.

A NORMAL, DIVERSE GUT MICROBIOTA IS CRITICAL TO HUMAN IMMUNITY AND CAN PREDICT INFECTION IN PATIENTS WITH AL

Infection is a common symptom in patients with newly diagnosed AL. In addition, the bone marrow suppression that occurs after chemotherapy in AL patients is also prone to infection. Infection is an important cause of death in patients with AL (40, 41). Gut microbiota is strongly associated with host.
immune function and resistance to infection. It is essential for the host to maintain a healthy immune system (42). It has been shown that gut microbiota can produce inhibitory products to suppress vancomycin-resistant enterococci (43). The use of carbapenem antibiotics in the empirical treatment of fever due to neutropenia in patients with AL reduces gut microbiota diversity and increases colonization by vancomycin-resistant enterococci (44). SCFAs, a metabolite of gut microbiota, promote the differentiation of T lymphocytes into effector T cells and regulatory T cells by inhibiting histone deacetylase (45). SCFAs also significantly inhibited the activity and infectivity of E. coli and protected intestinal epithelial cells from damage caused by C. difficile toxins (45, 46). Treatment of mice with broad-spectrum antibiotics resulted in a decrease in the number of hematopoietic stem cells and pluripotent progenitor cells in their bone marrow, as well as a decrease in granulocytes and B cells in the bone marrow and an increase in CD8+ T cells (47). This is the result of broad-spectrum antibiotics interfering with the STAT1 signaling pathway by consuming the gut microbiota (47). The aromatic amino acid metabolite production pathway of intestinal gut symbiont Clostridium sporogenes produces 12 compounds, 9 of which accumulate in host serum, indolepropionic acid being one of them. A decrease in indolepropionic acid increases the number of immune cells in mice, including neutrophils, monocytes, and memory T cells, and mice with decreased indolepropionic acid have greater intestinal permeability and are more likely to trigger an inflammatory response in mice (48). A normal and diverse gut microbiota is essential for human immunity and resistance to infection.

Causes of gut microbiota dysbiosis in AL patients associated with antibiotic use, nutrition, and chemotherapy (49). A study used a 16s rRNA-based analysis of bacterial flora in oral and fecal specimens from 97 AML patients every two weeks from induction chemotherapy to neutrophil recovery. They found that fecal flora diversity and high levels of Porphyromonadaceae at the time of induction chemotherapy were positively associated with a low risk of infection (50). In AML patients who developed infection during induction chemotherapy, baseline fecal flora diversity was significantly lower than in patients who did not develop infection, and flora diversity gradually decreased over the course of induction chemotherapy (51). In a study of ALL, chemotherapy was also found to have an impact on the diversity of the gut microbiota, with a significant decrease in the relative abundance of certain bacterial groups (e.g., Bacteroidetes) and a significant increase in the relative abundance of other groups (e.g., Clostridiae and Streptococcaceae) following chemotherapy (52). In contrast to the findings of some studies (44), this study concluded that antibiotic use does not alter the diversity of the gut microbiota (52). However, it has also been suggested that vancomycin use significantly reduces gut microbial diversity in patients with ALL, while no correlation was found with piperacillin tazobactam, antifungal drugs and other antibiotics (53). Some investigators concluded that the composition, but not the diversity, of the gut microbiome prior to chemotherapy was an independent predictor of febrile neutropenia during chemotherapy. The predominance of Aspergillus phylum and Enterococcus (relative abundance ≥30%) predicted febrile neutropenia; Enterococcus (relative abundance ≥30%) and Streptococcaceae predominate in predicting diarrheal disease (52). Intestinal nutrition deficiency can lead to loss of intestinal epithelial barrier function and increased susceptibility to infection (54).

In a study evaluating the effect of probiotics on chemotherapy-induced gastrointestinal side effects in patients with AL, patients with AL were randomized to an oral Lactobacillus rhamnosus group and a control group. The primary endpoint was the incidence of gastrointestinal side effects, with significant reductions in nausea, vomiting, and abdominal distention (P<0.05) and a relative risk of diarrhea (0.5) in the oral probiotics group (55). Patients in the control group of this study did not take placebo and subjective error may occur. In a study of gut microbiota recovery in mice and humans, it was found that neither probiotics nor spontaneous recovery restored gut microbiota diversity after 4 weeks of disruption cessation with antibiotics. Importantly, probiotics significantly delayed recovery to baseline microbiome richness at all time points tested, compared to spontaneous recovery (56). However, gut microbiota diversity was indistinguishable to control after 8 days of fecal transplantation (56). The use of probiotics can only restore one or a few specific gut microbes, but not the function of a complex microbial community. Therefore, supplementation with one probiotic alone may have adverse effects on the host (49), or may even result in probiotic-induced infections. A patient who received autologous hematopoietic stem cell transplantation took probiotic-rich yogurt due to severe diarrhea, and a week later the patient developed infectious shock (57). Strain-specific PCR analysis showed that the pathogen was Lactobacillus rhamnosus GG, which was consistent with the probiotics in the yogurt (57). Another child with Philadelphia chromosome-positive ALL underwent chemotherapy and developed neutropenic fever with blood cultures suggestive of Bifidobacterium breve, which may be pathogenic when host immunity is reduced (58). In patients with AL, the diversity and composition of gut microbes correlate with whether or not an infection occurs or what type of infection occurs, and they are influenced by factors such as chemotherapy and antibiotics. Probiotics can reduce gastrointestinal symptoms such as diarrhea, but opportunistic infections with probiotics have also been reported in the presence of neutrophil deficiency.

The gut microbiota is closely related to the normal immunity of the organism (42). Among the infections present in patients with AL, the gut microbiota has been associated with diarrhea, neutropenic fever, and sepsis, and its diversity and composition are predictive of infection in patients (45–52, 59).

**GUT MICROBIOTA AFFECTS A WIDE RANGE OF DRUGS USED TO TREAT AL**

There is increasing evidence that host responses to chemotherapeutic agents can be influenced by the gut...
microbiota, primarily in terms of promoting drug efficacy; abrogation of anticancer effects; and modulation of toxicity (60). Gut microbiota regulates chemotherapeutic drugs through key mechanisms such as reduced diversity, ecological variation, translocation, immune regulation, metabolism, enzymatic degradation (60).

Cyclophosphamide (CTX) is commonly used to treat ALL, lymphoma, and other tumors. Vlua S et al. investigated the role of gut microbiota in CTX therapy through a mouse model (61). Non-myeloablative doses of CTX and adriamycin treated mice resulted in shortened small intestinal villi, disrupted epithelial barrier, and interstitial edema 48 hours after drug application (61). After the disruption of the intestinal barrier, some intestinal microorganisms are transferred to the mesenteric lymph nodes and spleen, and intestinal microorganisms including Lactobacillus johnsonii, Lactobacillus murinus and Enterococcus hirae can be cultured in these lymphoid organs (61). CTX treatment induces selective translocation of gut microbiota, for example, leading to reduced abundance of Lactobacillus and Enterococcus in the small intestine of mice. Some intestinal bacteria (Lactobacillus johnsonii, Lactobacillus murinus and Enterococcus hirae) stimulate the production of specific “pathogenic” T helper 17 (pT(H)17) cells. Killing these bacteria with antibiotics reduced the pT(H)17 response in tumor-bearing mice and made them resistant to CTX. Adaptive transplantation of pT(H)17 cells could recover the anti-neoplastic efficacy of CTX in part. In another study, calorie restriction was found to significantly enhance the gut microbiota with Lactobacillus and Trichinella, which are shown to relieve inflammation and improve intestinal barrier function. Calorie restriction reshapes stronger gut microbiota and may help reduce side effects of CTX (62).

Methotrexate (MTX) has the function of anticancer and immunosuppressive, and is one of the main drugs used in the treatment of ALL, central nervous system leukemia, and rheumatic immune system diseases (63–65). Common side effects of MTX include mucositis and intestinal damage (66, 67). MTX has significant effects on the amounts, diversity and major components of the mouse gut microbiota (68). MTX-treated mice showed reduced intestinal microbial diversity and changes in the composition of the gut microbiota, such as a dramatic decrease in the number of ruminal cocci and a dramatic increase in the number of members of the Lachnospiraceae (68). B. fragilis was significantly reduced by MTX treatment, and there was a tendency for the proliferation rate of B. fragilis to decrease with increasing M1 macrophage density (68). The mucosal muscle lesions seen in mttx-treated mice were significantly relieved by gavage with fragile B (68). Fragile B. markedly attenuated the MTX-induced increase in M1 macrophages, and the decrease in fragile B. in turn aggravated the MTX-induced macrophage imbalance, resulting in positive feedback, which led to intestinal tissue injury (68). Chronic long-term MTX exposure induced changes in the gut microbiota composition and function, with the relative abundance of the Firmicutes being higher than that of the Bacteroidetes at low doses of MTX, but the opposite at high doses of MTX. The relative abundance of the Firmicutes was positively correlated with the 48-h fecal excretion of 2,4-diamino-N-10-methylpteroid acid (DAMPA), a metabolite of MTX. Long-term exposure to MTX can alter the composition and function of the microbiome, which in turn affects its ability to detoxify MTX (69). MTX largely alters the human gut microbiota. Changes in gut flora and gene abundance were different between patients who responded to MTX and those who did not. MTX affects a variety of conserved human gut bacterial pathways, leading to reduced host immune activation (70).

PD-1 is a kind of immune checkpoint, PD-1 binding with PD-L1 will inhibit T cells and block the anti-tumor immune response. Anti-PD1 has been approved by FDA for the treatment of melanoma, non-small cell lung cancer, renal cell carcinoma and other tumors (71). In a mouse model, enhanced PD-1 expression on circulating CD8 T cells leads to the progression of AML (72). A phase II clinical trial of azacitidine combined with nivolumab in patients with relapsed/refractory AML obtained an encouraging result, with an overall response rate (ORR) of 33% (73). PD-1 inhibitors have promising applications in the treatment of AL, and the gut microbiota also interacts with PD-1 inhibitors. In a mouse model study of epithelial tumors, primary resistance of tumors to immune checkpoint inhibitors (ICIs) was found to be attributable to unusual composition of the gut microbiome, and the clinical benefit of ICIs in patients with late-stage cancer could be inhibited by antibiotics (74). Germ-free or antibiotic-treated mice receiving fecal microbiota transplantation (FMT) from patients with tumors that responded to ICIs improved the antitumor effects of PD-1 blockade, but FMT from patients with tumors that did not respond to ICIs had no such effect (74). Metagenomics of patient fecal samples revealed a correlation between the relative abundance of Akkermansia muciniphila and patient response to ICIs. Administration of oral supplementation of A. muciniphila to mice unresponsive to FMT restored the efficacy of PD-1 blockade in an interleukin-12-dependent manner (74). In another study of melanoma patients, researchers found that the effect of anti-pd-1 immunotherapy was associated with greater abundance of Bifidobacterium longum, Collinsella aerofaciens, and Enterococcus faecium in the patients’ stools (75). Transplanting feces from patients who responded to anti-pd-1 into germ-free tumor-bearing mice enhanced the T-cell effect and improved tumor control (75). In a study of the gut microbiota of melanoma patients receiving PD-1 immunotherapy, investigators found that the diversity of the flora and the relative abundance of Ruminococcaceae family were significantly higher in patients who responded to PD-1 immunotherapy (76). In immunotherapy for liver cancer, researchers also found more taxa richness and higher gene counts in the stools of patients who responded to anti-PD-1 (77). Gut microbiota correlates with the efficacy of ICIs in a wide range of tumors, with higher diversity of gut microbes associated with better efficacy. Whether the efficacy of anti-PD-1 therapy in patients with AL is also associated with the gut microbiota needs to be urgently investigated.
There are no studies on the interaction of gut microbiota with drugs used to treat AML (e.g., erythromycin, cytarabine, etc.), which is a future direction to investigate the relevance of gut microbiota to AL therapeutic drugs.

**THE PLACE OF GUT MICROBIOTA IN ALLO-HSCT: FOR BETTER OR FOR WORSE?**

Allo-HSCT is currently one of the most effective treatments for AL (78). Allo-HSCT means that hematopoietic stem cells collected from a healthy donor are infused into a patient with leukemia, and the donor’s immune cells can attack the leukemia cells to produce a graft-versus-leukemia (GVL) effect (79). The main side effect of allo-HSCT is graft-versus-host disease (GVHD) (79, 80). GVHD means that the immune cells of the donor attack the normal tissues of the patient, and the common organs are the skin, liver, and gastrointestinal tract, and the patient may present with diarrhea, vomiting, abdominal pain, jaundice, sclerosis of the skin, and other harmful manifestations (79). Acute graft-versus-host disease (aGVHD) occurs in 30-50% of all allo-HSCT patients, and chronic graft-versus-host disease (cGVHD) occurs in 35-50% of patients (81, 82). Prevention of GVHD includes the use of cyclosporine, tacrolimus, mycophenolate mofetil (MMF), methotrexate, etc (83). Despite the use of these drugs, many patients still develop GVHD, and glucocorticoids are an effective treatment for both aGVHD and cGVHD (83). But glucocorticoids can cause side effects such as osteoporosis, susceptibility to infection, increased blood sugar and blood pressure, necrosis of femoral head and so on. In summary, GVHD affects the quality of survival, mortality in allo-HSCT patients. Gut microbiota plays an important role in the development and treatment of GVHD (84).

Gut microbiota correlates with the development of GVHD. In 1974, van Bekkum DW et al. found that GVHD occurred in mice treated with lethal total body irradiation and allo-HSCT, with a mortality rate of 95% within 100 days, whereas mice treated similarly in a sterile state or given a colonization-resistant flora had almost no GVHD (85). This study confirmed in an animal model that the microbiome is associated with GVHD. In a study of mice, it was found that the gut microbiota diversity of mice with GVHD decreased significantly during the first 2 weeks of bone marrow transplantation, and in terms of changes in bacterial subgroups, the number of Lactobacillus increased significantly in the ileum, while the number of Clostridium and other Firmicutes decreased significantly (86). The authors also found that removal of Lactobacillus prior to bone marrow transplantation aggravated GVHD in mice and that reintroduction of a dominant species of Lactobacillus mediated protection against GVHD (86).

The presence of disruption of the gut microbiota early in bone marrow transplantation is also a potential risk factor for GVHD (86). We already know that SCFAs can reduce DNA damage in blood and gastrointestinal tissues during radiation damage (31), and SCFAs inhibit histone deacetylation (45). SCFAs mainly include formate, acetate, propionate and butyrate (87). Mathewson ND et al. analyzed SCFAs produced by the microbiome of allo-HSCT mice and found that only butyrate was significantly reduced in the intestinal tissues of transplanted recipients, resulting in reduced levels of histone deacetylation in small intestinal epithelial cells. In contrast, increasing the level of butyrate in the small intestine restored histone deacetylation, protected small intestinal epithelial cells, and reduced the severity of GVHD (88). The mechanism by which butyrate protects small intestinal epithelial cells may be through the regulation of histone deacetylation to increase anti-apoptotic genes. In butyrate-treated small intestinal epithelial cells, Bax and Bak were significantly reduced and the anti-apoptotic protein BCL-B was significantly increased (88). In the study of SCFAs and cGVHD, higher SCFAs (mainly butyrate and propionate) in the plasma of patients at 100 days post-transplantation were associated with decreased cGVHD (89). Golob JL et al. prospectively collected 669 stool samples from 66 allo-HSCT patients, once a week from pre-transplant to 100 days post-transplant. Their analysis revealed that the presence of Lachnospiraceae in stool was negatively associated with GVHD when neutrophils recovered after transplantation, whereas the presence of oral Actinobacteria and oral Firmicutes was positively associated with GVHD (90). A study by Han L et al. found more gut microbiota diversity in the non-aGVHD group than in the aGVHD group (91), they evaluated the gut microbiota to predict GVHD and found that Lachnospiraceae and Peptostreptococcaceae were negatively associated with GVHD, while Enterobacteriaceae were positively associated with aGVHD (91). Lachnospiraceae was negatively correlated with severe aGVHD, and levels of all major SFCAs were significantly decreased in patients with severe aGVHD, with a 75.8% decrease in acetate, a 95.8% decrease in propionate, and a 94.6% decrease in butyrate, while propionate levels were retained in patients with mild aGVHD. Butyrate was severely reduced in all gastrointestinal stages of aGVHD (92). Patients with ≤10% Lachnospiraceae in stool specimens at transplant pretreatment were significant contributors to overall mortality, and reduced gut microbiota diversity at day 10 post-transplant significantly increased the risk of GVHD (93). These studies all confirmed that abundant Lachnospiraceae and intact gut microbiota diversity were negatively correlated with GVHD (90–93). Lachnospiraceae-dominant gut microbiota mice also can withstand high doses of radiation (31), Lachnospiraceae and SCFA (propionate) are used to modulate host resistance to high doses of radiation by promoting hematopoiesis and gastrointestinal recovery (31).

Another product of the gut microbiota, indole, has also been shown to improve GVHD. Indole is produced by certain intestinal microflora (e.g., Escherichia coli, Lactobacillus, Bacteroides) (94). Indole-3-carbaldehyde (ICA) is an exogenous indole derivative. ICA upregulated genes associated with type I interferon (IFN1) responses, limited intestinal epithelial damage, reduced inflammatory cytokine production, and decreased GVHD pathology and GVHD mortality without affecting donor T cell-
mediated graft-versus-leukemia responses (95). Although SCFAs and indoles, metabolites of the gut microbiota, have been shown to reduce the severity of GVHD, another circulating gut microbial metabolite, TMA N-oxide (TMAO), has been reported to enhance M1 macrophage polarization, and polarized inflammatory macrophages establish an environment for Th helper type 1 (Th1) and Th17 responses, thereby increasing GVHD (96).

The use of broad-spectrum antibiotics can increase the severity of GVHD. In a mouse model of allo-HSCT, treatment with imipenem-cilastatin was found to disrupt intestinal microbiota and increase the severity of GVHD (mainly increased GVHD pathology in the colon) in mice with GVHD, resulting in increased mortality (97). However, if the mice did not develop GVHD, treatment with imipenem-cilastatin did not affect the survival of the mice (97). Why did the use of imipenem-cilastatin lead to increased severity of GVHD in mice? Further studies revealed that in these mice, donor effector CD4+ T cells (CD25+FoxP3-) and interleukin-23 (IL-23) were increased and granulocytes were infiltrated in their colon (97). AL patients treated with intensive chemotherapy and broad-spectrum antibiotics destroy the diversity of the intestinal microbiota (98). The use of autologous fecal microbiota transfer (AFMT) restores intestinal flora diversity and is safe for immunocompromised patients (98). Whether AFMT can improve GVHD requires further study.

Gut microbiota correlates with the prognosis of allo-HSCT. Kuwasakabe S et al. studied the gut microbiota of normal, autologous HSCT patients and allo-HSCT patients, and they found that in allo-HSCT patients with skewed microbiota had a higher frequency of more complications and death after transplantation than the subgroup with preserved microbiota composition (93). The diversity of the gut microbiota is an independent predictor of mortality in allo-HSCT patients, Taur Y collected stools from 80 allo-HSCT patients for analysis, and the percentage of patients maintaining high, intermediate, and low diversity of the gut microbiota was 26 (32.5%), 20 (25.0%), and 34 (42.5%), and their 3-year overall survival rates were 67%, 60% and 36%, respectively (99). Peled JU et al. analyzed 8767 stool samples from 1362 patients who received allo-HSCT from a total of four transplant centers in the United States, Germany, and Japan and found that higher gut microbiota diversity at the time of neutrophil engraftment was associated with lower patient mortality, patients with low gut microbiota diversity were positively associated with transplant-related mortality and a high risk of GVHD (100). A study on gut microbiota and pulmonary complications after allo-HSCT found that low gut microbiota diversity was associated with PCs occurring pre-engraftment, whereas γ-proteobacteria dominant gut flora was associated with PCs post-engraftment (101). PCs post-engraftment and γ-proteobacteria dominant gut flora were associated with mortality (101). In a study of a total of four bone marrow transplant centers from the United States, Japan, and Germany, the investigators defined Enterococcal dominance as having a relative genic abundance of ≥0.3 in any fecal sample (102). The incidence of Enterococcus dominance was high at all centers, with 441 of 1101 patients (40.1%) at Memorial Sloan Kettering Cancer Center (MSKCC) and 103 of 224 patients (46.0%) at the other 3 centers having E. faecium dominance between day -20 and day +80 of allo-HSCT (102). Early post-transplant (day 0 to +21) Enterococcus dominance significantly reduced overall survival and increased GVHD-related mortality, increasing the risk of severe aGVHD (102). In a mouse model of allo-HSCT, an increase in Enterococcus in the feces of mice was also positively associated with GVHD, and the absence of lactose in the diet significantly reduced the colonization of Enterococcus after transplantation and attenuated GVHD (102). Lactose-free diets may be used in clinical settings to attenuate the growth of pathogens such as enterococci and improve clinical outcomes.

### TABLE 1 | Mechanisms by which the gut microbiota affects GVHD.

| Influence of gut microbiota on GVHD | Mechanisms | References |
|-------------------------------------|------------|------------|
| Lactobacillus shows improvement in GVHD | Butyrate improved IEC junctional integrity, decreased apoptosis | (88, 99) |
| Disruption of gut microbiota diversity aggravates GVHD | – | (86, 91, 93) |
| Butyrate, a metabolite of the gut microbiota, attenuates GVHD | – | (90-93) |
| Lachnospiraceae is negatively associated with GVHD | Indole upregulates genes associated with IFN1 response, limits intestinal epithelial damage, reduces inflammatory cytokine production, and decreases GVHD pathology and GVHD mortality. | (95) |
| Indole, a metabolite of the gut microbiota, improves GVHD GVHD progression was induced by TMAO, a circulating gut microbial metabolite | TMAO enhances M1 macrophage polarization and establishes an environment for Th1 and Th17 responses | (96) |
| Antibiotic use alters the composition of the gut microbiota and increases GVHD | Loss of protective colonic mucosa and impairment of intestinal barrier function due to antibiotics | (97) |
| Obesity aggravates GVHD by affecting the gut microbiota | Obesity reduces the diversity of the gut microbiota and reduces Clostridiaceae abundance, leading to increased intestinal permeability, transintestinal transit of endotoxins, and radiation-induced gastrointestinal damage in mice | (107) |
| The microbiota controls MHC-II at the pre-transplant IEC | Microbiota depletion inhibits IL-12/23p40 production by ileal macrophages, IL-12/23p40 prevents upregulation of MHC class II cells on IECs and initiation of lethal GVHD in the gastrointestinal tract | (108) |
| The gut microbiota metabolite sensor G-protein-coupled receptor 43 (GPR43) attenuates gastrointestinal GVHD | GVHD protection by SCFAs requires GPR43-mediated ERK phosphorylation and activation of NLRP3 inflammasome in host non-hematopoietic target tissues | (109) |
Interestingly, it has been shown that the relative abundance of Eubacterium hallii is higher in minimal residual disease (MRD)-negative multiple myeloma patients compared to MRD-positive patients (103). Whether the relative abundance of gut microbiota is associated with risk stratification of AL and MRD has not been studied, which warrants further research.

FMT is a treatment for multidrug-resistant bacteria colonization and GVHD in patients undergoing allo-HSCT. Battipaglia G et al. performed FMT in 10 patients colonized by multidrug resistance; eight of these patients were colonized by carbapenemase-producing bacteria and two by vancomycin-resistant enterococci; four of them underwent FMT before allo-HSCT and the other six underwent FMT after allo-HSCT; seven of the 10 patients achieved decolonization (104). We already know that disruption of gut microbiota diversity can exacerbate GVHD, but patients are in a state of severe immunosuppression and the exact mechanism by which it reduces recurrence needs to be confirmed by further studies.

In general, the gut microbiota may play an important role in the pathogenesis, progression and treatment of AL by influencing human immunity. The gut microbiota has been found to resist damage from radiation and benzene, which may cause AL. The gut microbiota is also closely related to the body’s normal immunity, and in AL, the diversity of the gut microbiota is associated with multiple infections. When AL patients undergo chemotherapy, the gut microbiota is affected and, in turn, the gut microbiota affects the efficacy of multiple chemotherapeutic agents. The diversity of the gut microbiota and its metabolites are strongly associated with the occurrence of GVHD, mortality and relapse of allo-HSCT. More studies can be conducted in the future to focus on the relationship between the gut microbiota and AL, especially that the gut microbiota improves the treatment outcome and prognosis of AL patients by regulating human immunity.

AUTHOR CONTRIBUTIONS

TM and YC wrote the manuscript. L-JL and L-SZ wrote and critically revised the manuscript. All authors contributed to the article and approved the submitted version.

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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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