Rheumatoid arthritis (RA) is a disabling autoimmune disease with invasive arthritis as the main manifestation and synovitis as the basic pathological change, which can cause progressive destruction of articular cartilage and bone, ultimately leading to joint deformity and loss of function. Since its introduction in the 1980s and its widespread use in the treatment of RA, low-dose methotrexate (MTX) therapy has dramatically changed the course and outcome of RA treatment. The clinical use of this drug will be more rational with a better understanding of the pharmacology, anti-inflammatory mechanisms of action and adverse reaction about it. At present, the current clinical status of newly diagnosed RA is that MTX is initiated first regardless of the patients’ suitability. But up to 50% of patients could not reach adequate clinical efficacy or have severe adverse events. Prior to drug initiation, a prognostic tool for treatment response is lacking, which is thought to be the most important cause of the situation. A growing body of studies have shown that differences in microbial metagenomes (including bacterial strains, genes, enzymes, proteins and/or metabolites) in the gastrointestinal tract of RA patients may at least partially determine their bioavailability and/or subsequent response to MTX. Based on this, some researchers established a random forest model to predict whether different RA patients (with different gut microbiome) would respond to MTX. Of course, MTX, in turn, alters the gut microbiome in a dose-dependent manner. The interaction between drugs and microorganisms is called pharmacomicrobiology. Then, the concept of precision medicine has been raised. In this view, we summarize the characteristics and anti-inflammatory mechanisms of MTX and highlight the interaction between gut microbiome and MTX aiming to find the optimal treatment for patients according to individual differences and discuss the application and prospect of precision medicine.

Keywords: methotrexate (MTX), rheumatoid arthritis (RA), gut microbiome, pharmacomicrobiology, precision medicine
1 INTRODUCTION

Rheumatoid arthritis (RA) is a systemic autoimmune disease characterized by persistent synovitis, chronic and aggressive arthritis. Although the treatment of RA has made great progress in recent years, the overall remission rate of RA is still not optimistic, and it is almost impossible to reverse once bone or joint destruction occurs (1, 2). The pathogenesis of RA is still not fully understood, which may be caused by the complex interaction between genetic environment and immune factors, resulting in immune system dysregulation and lack of autoimmune tolerance. Although several effective treatments have been developed recently, low-dose methotrexate (MTX) remains the anchor agent for RA treatment, which is generally considered to enhance the efficacy of most biologics in RA (3, 4). However, a large proportion of RA patients do not respond to oral MTX or have severe adverse reactions, especially gastrointestinal toxicity, which is common, which can be troublesome (3–6). Therefore, it is urgent to find out the possible reasons for individual differences and the biological indicators that can predict the clinical efficacy of MTX to guide clinical medication.

An adult human has about 2×10^14 host cells, but this is only a tenth of the bacteria that live in the mucosae of the body (the intestine, reproductive and respiratory tract) and outside the body (hair and skin). Thus, the number of microbes living in the human body is extremely huge (7). These microbial communities are intimately involved in our body’s metabolism, and some emerging research is trying to deciphering the complex cross-border communication network between our immune system and the microbial community that exists in our bodies (8). The gastrointestinal tract is home to about 1,500 species of more than 100 trillion microbes, with the highest diversity and density of microbes ever seen (8). The microbiome is involved in many physiological processes, including digestion and metabolism, and the development and regulation of the immune system. In turn, the host provides living space and nutrients for the microbes. Host-microbial interaction, especially in the gut, promotes the development and regulation of the host immune system and is essential for the stability of the entire complex microbial community (9, 10). In other word, gut microbiome is an important immune organ and plays an indispensable role in immune response and tolerance (11, 12). The gut microbiome is increasingly considered to be involved in the processing of various exogenous substances and can also influence host responses to various compounds, including all kinds of drugs. And in this process, gut microbes are evolving under the influence of exogenous compounds. Metagenomics of intestinal microorganisms plays an important role in the efficacy and toxicity of many drugs, and its changes may affect the performance of drugs (13–16).

Now the most recent view on MTX and the gut microbiota is that gut microbiome can affect the bioavailability of MTX and some characteristics of gut microbiome can be considered as a predictor of clinical response of this drug (17). In addition, gut microbiome is associated with adverse reactions to MTX, especially gastrointestinal reactions (18). Of course, MTX also affects the amount, diversity, and major components of the gastrointestinal microbiota (18, 19). Pharmacomicrobiology, based on drug-microbe potentially complex interactions has been used in various fields (20, 21). Precision medicine based on pharmacomicrobiology, while strictly integrating clinical factors and host genomics will be the development direction of rheumatoid diseases including RA in the near future (17, 21). In this review, we focus on the pharmacology, anti-inflammatory mechanism, adverse reactions of MTX and diagnostic and predictive effect of gut microbiome on clinical efficacy of oral MTX to recognize the role of pretreatment of gut microbiome in improving clinical efficacy of MTX, in order to provide a new therapy for the targeted treatment of RA patients who fail to respond within the early therapeutic window of opportunity.

2 MTX AND ITS MECHANISMS OF ACTION IN RA

MTX (4-amino-4-deoxy-N-10-methylpteroylglutamic acid) is an anti-folate cellular immunosuppressant. Low doses (<15mg/week) of MTX are widely used to treat autoimmune diseases such as RA or psoriasis (19, 22). However, higher doses of MTX (typically between 15 to 500 mg/kg) are effective in the treatment of cancer diseases, mainly for acute lymphoblastic leukemia, and osteosarcoma and lymphoma (19, 23–25). Ever since Sidney Faber discovered MTX in the 1980s, low-dose MTX is generally the first-line drug for the treatment of RA (26, 27). It is possible that MTX has been enduring because it is well tolerated, safe, and significantly less costly (28–30). It was a long and difficult journey from the initial discovery of MTX to widespread use around the world (26) (see Figure 1). Currently, once RA is diagnosed, rheumatologists recommend taking MTX orally. If the patient has a poor respond or strong adverse reaction to MTX, the patient is advised to continue taking it along with the biologic, since MTX appears to have an additive effect on biologic drugs (4). At low doses, MTX exhibits anti-inflammatory properties, but at high doses or for long periods of time, it may cause the pathological multi-organ toxicity, including gastrointestinal, myelotoxicity, cardiotoxicity, nephrotoxicity, and hepatotoxicity, which limits its therapeutic potential (18, 31–33). Why do different individuals respond differently to MTX? As we learn more about the pharmacological effects, anti-inflammatory mechanisms and interactions with gut microbes of MTX, the answer to this question may become more and more clear. In fact, emerging data suggest that it is possible to identify appropriate biomarkers of the gut microbiome to determine who responds best to MTX and has the fewest side effects – that is, who will benefit the most from MTX treatment.

2.1 Pharmacokinetics of MTX and Gut Microbiome

MTX therapy for RA is usually effective in the range of 15 to 25 mg and is usually initiated as monotherapy (22, 27). The intestinal tract has a limited ability to absorb MTX, so oral it has limited bioavailability in vivo (34, 35), which may be related to gut microbiota. The maximal absorption of a single oral dose
is <25mg (3). Therefore, rheumatologists recommend that patients take the required dose of medication orally for one day to obtain better clinical response from MTX, rather than taking a daily dose. Due to the complex gastrointestinal toxicity of MTX, attention has shifted from oral to subcutaneous. A recent study (36) retrospectively analyzed for pharmacokinetics clinic attendances at a large North-East England hospital and concluded when the dose of MTX in RA patients > 15 mg/week, subcutaneous method is safer than oral medicine. Specifically, the subcutaneous approach reduced the incidence of neutropenia and gastrointestinal symptoms. One problem, however, is that patients who experience adverse events with oral MTX are less likely to switch to subcutaneous treatment and are likely to experience the side effects of oral MTX again. Therefore, the search for predictors of MTX before treatment is irreplaceable.

After taking MTX orally, it is first actively absorbed by the proximal jejunum. The degree and dose of absorption are attributable to saturation of reduced folate carrier 1 (RFC1) (37). The absorptivity of MTX is so high that the plasma concentration reached its maximum value within 0.75 ~ 2 h after oral Administration (26). But this drug has a relatively short half-life of about 1 hour (26) and after 18 hours its concentration in the serum is almost zero and undetectable (3). MTX enters the circulation from the gut, where 35 to 50% of the drug binds to albumin (26). Most MTX is transferred by RFC1 into cells, mainly red blood cells, white blood cells, synovial cells and liver cells. In red blood cells, MTX is polyglutamated by adding glutamate groups to the adjacent glutamate leaf terminal carboxyl group in the γ linkage (3, 26, 27). And the whole reaction is catalyzed by the enzyme folic acid-glutamate synthetase. Red blood cells are a repository for MTX. Thus, the toxicity of MTX can also be affected by the low dose of weekly administration, which should be considered in clinical use of this drug. In the liver, MTX is oxidized to 7-hydroxyl MTX (7-OH-MTX) by aldehyde oxidase (AO), and then both parent compounds and metabolites mixed into the bile which then passes through the common bile duct and into the intestine, to duodenum, to be precise (26, 38, 39). Here, they can undergo further metabolism and biotransformation through the gut microbiome (19, 40, 41). Drugs and metabolites that enter the intestinal tract can return to the liver via the portal vein, which is known as the enterohepatic circulation. Cholates (unbound bile salts) can inhibit this circulation, which is very important to mitigate potential adverse events and has been showed to reduce the potential toxicity of MTX (42, 43). MTX and its metabolites are mainly excreted in urine (26, 27, 44), up to 30% are metabolized in bile (26, 45) however, only 1-2% are excreted in feces (26, 35).

Gut microbiome can directly affect the bioavailability of MTX (19). Glutamate carboxypeptidase 2 (CPDG2) is an enzyme found in many gut bacteria. The terminal glutamate residues of MTX and 7-OH-MTX can be cleaved to generate 2, 4-diamino-10-methylpteroidic acid (DAMPA) and 7-hydroxy-DAMPA metabolites with the assistance of CPDG2, respectively (41, 46, 47). In addition, other bacterial enzymes such as p-aminobenzoyl-glutamate hydrolase, found in E. coli, have also been shown to catalyze this reaction (48) (see Figure 2). The hydrolysis of glutamate by CPDG2 is considered to be a detoxification process. Furthermore, CPDG2 has been approved as a palliative for cancer patients with delayed MTX clearance and acute renal toxicity (24). Several clinical studies have also reported the usefulness of CPDG2 as a rescue drug (45, 50). This is just one example of how the gut microbiome influences MTX metabolism, and there are certainly other, deeper connections between MTX and gut microbes that we need to explore.

### 2.2 Anti-Inflammatory Mechanisms of MTX

There is growing evidence that many of the mechanisms by which MTX inhibits the inflammatory response have been
hypothesized. In this section, we focused on exploring the anti-inflammatory mechanism of MTX to guide rational clinical drug use.

MTX and folates. MTX reaction products inhibit dihydrofolate reductase (DHFR) and affect folic acid metabolism, thus restraining purine and pyrimidine synthesis and downstream DNA synthesis (3, 26, 27, 51). The reduction of purine and pyrimidine synthesis suppresses the T lymphocytes, a key cell in the immune response, so MTX has anti-inflammatory effect (26, 52). However, this drug as a folate analogue can cause a decrease in peripheral blood white blood cell count which is still considered to be a toxic reaction in treating inflammatory diseases (3), although the dose of MTX used to treat RA is 100 to 1000 times lower than that used to treat malignancies. It is now common sense that taking folic acid (except on MTX days) can reduce or even prevent the toxic effects of this immunosuppressant in patients (53–55).

Given that even supplementation with folic acid during medication can prevent a decrease in peripheral white blood cell count and that MTX can still inhibit progression in patients with rheumatic disease, the anti-inflammatory effect of MTX may not be as necessary to inhibit cell proliferation (3).

MTX and adenosine. There is growing evidence that MTX causes the release of adenosine, a powerful stimulator of adenosine receptors (including A1a, A2a, A2b and A3) that inhibits nearly all types of inflammatory cells (56). Studies in mice demonstrated that MTX increased the release of adenosine in inflammatory tissues (57, 58). According to a randomized investigational study (59), ingestion of non-selective adenosine receptor antagonist such as caffeine can reduce the body’s response to MTX therapy, which confirmed MTX treatment can increase the body’s really adenosine release. Recently, emerging evidence suggests that the mechanism by which regulatory T cells (Treg, a type of negatively regulated immune cells) reduce the degree of immune response and thus inhibit inflammation may be related to the production and release of adenosine (56, 60). Treg cells produce adenosine through dephosphorylation of ATP, a process mediated by CD39 and CD73 (61–63). If CD39 or CD73 is relatively low on Treg cell level, such as in RA, this process is impeded, resulting in a poor clinical MTX response (64).

MTX and reactive oxygen species (ROS). Nitric oxide synthase (NOS) is an isoenzyme found in endothelial cells, macrophages, and nerve cells, which normally promotes the production of NO. Tetrahydrobiopterin (BH4) is a cofactor and ligand of endothelial nitric oxide synthase (eNOS). In the absence of BH4, NOS does not catalyze the production of NO, but instead produces ROS, such as hydrogen peroxide (a damage factor for human tissues and organs), a process known as nitric oxide synthase uncoupling (65–67). It has been proved that MTX can inhibit nitric oxide synthase uncoupling (3). MTX suppresses the reduction of dihydrofolate and dihydrobiopterin (BH2), that is, increases their concentration, which in turn increases the concentration of BH4 [BH2 is restored to BH4 by dihydrofolate reductase (DHFR)], thereby reducing the production of ROS.

MTX and cytokines. MTX alters the cytokine profile, specifically inhibiting the production of pro-inflammatory cytokines (27, 68, 69). It has been shown that T cells isolated and activated from RA patients treated with MTX have a reduced ability to produce inflammatory cytokines such as IFN-γ, IL-4, IL-3, and TNF (70). MTX treatment also reduced the number of TNF-positive CD4 T cells and increased the number of IL-10 CD4 T cells (71). Which is beneficial for RA patients to manage their disease.

Other anti-inflammatory mechanisms have also been reported, including inhibition of transmethylation (3), regulation of the expression of some long non-coding RNAs (3), and reduction of
chemotaxis and adhesion of inflammatory cells (27) (see Figure 3). Considering the important efficacy, wide application and numerous adverse reactions of MTX, further research on the mechanism of action of MTX is to optimize treatment.

### 2.3 Toxic Effects of MTX

In the nearly 40 years of MTX use, an increasing number of toxic effects have been observed, including gastrointestinal, myelotoxicity, cardiotoxicity, nephrotoxicity, and hepatotoxicity. Currently, enterotoxicity is the most common side effect and the main factor limiting its administration, mainly manifested as MTX-induced intestinal mucosal inflammation, which is now increasingly evidenced to be related to gut microbiome (18). It has been suggested in the past that the toxic effects of long-term oral low doses of MTX are inextricably related to the antifolate properties of the drug (3). So it has become common knowledge to take folic acid along with MTX to prevent its toxic effects. Now, the latest international view suggests that the mechanism of folic acid supplementation to reduce the toxic effect may be related to intestinal flora, especially the probiotics such as Bacteroides and Bifidobacterium. And some researchers think that some of the toxic effects of MTX may be caused by MTX-mediated adenosine release. MTX promotes the release of adenosine in the inflammatory sites, which binds to receptors in the brain to suppress the inflammatory response and causes drowsiness (72–74). So many patients feel tired on the day they take MTX and other patients developed rheumatoid nodules (75) and liver fibrosis (76, 77).

Another serious and unpredictable side effect is pulmonary toxicity, which occurred in 1% of the 500,000 RA patients treated with MTX (78, 79), mainly including interstitial pneumonia and pulmonary fibrosis (80). The mechanisms include genotype mutation and transport inhibition, and are related to the p38 Mitogen-activated protein kinase (MAPK) pathway and interleukin-8 (IL-8) (81). These mechanisms can be used to optimize drug therapy, such as serratiopeptidase (PSTD) combined with fisetin (FST) against MTX-induced pulmonary toxicity (82). One thing to be aware of is hypersensitive pneumonia (HP), a diffuse interstitial lung disease caused by sensitization of susceptible individuals to repeated and prolonged exposure to large amounts of antigen (83). It is often associated with exposure to external allergens (such as poultry dust, mould and tobacco) and medications including cytotoxins (such as MTX) and antiepileptic drugs (such as carbamazepine) (84). Some HP patients may recover, while others develop pulmonary fibrosis, which may be related to the microbiota of the lower respiratory tract (85). For example, Proteobacteria are the predominant in HP, while Firmicutes dominate in pulmonary fibrosis.

However, a recent randomized, double-blind, placebo-controlled trial (86) showed that low doses of MTX reduced the incidence of complications from cardiovascular events in patients with RA and severe cardiovascular risk. Another new and particularly worrisome event is skin cancer (86). Various European biologics registers found a higher incidence of skin cancer in RA patients taking MTX than in the general population (87–89). This has been attributed to long-term immunosuppression, and these new data support this. Long-term oral low-dose MTX, on the one hand, slows disease progression and reduces the risk of subsequent complications such as lymphoma and cardiovascular events, and also increases the incidence of skin cancer.

### 2.4 Therapeutic Implications of MTX

Since the discovery of rheumatoid arthritis in the middle of the last century, MTX is still the gold standard for RA, despite the recent emergence of various drugs. Factors that can predict clinic response and prognosis to the drug are especially important to reduce disease activity and unnecessary potential toxic effects. Lately, a machine learning analysis of whole blood transcriptome data from RA patients (90) holds that gene expression of multiple inflammatory pathways may have predictive value for clinical response to MTX, and that gene expression of anti-I interferon response is the most valuable predictor. However, the results were not satisfactory, and the whole genes containing all

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**FIGURE 3** | Anti-inflammatory mechanisms of MTX. MTX has many anti-inflammatory mechanisms. Some of these pathways are known, and there are certainly many more that need to be explored.
the gene expression had higher predictive value than the individual interferon gene alone (3, 90). Meanwhile, several other studies have sought to identify clinical and/or laboratory markers that might predict clinical responses to MTX, but all ended in failure (91–93). Despite high hopes for genetic markers that identify MTX responses, to date, no factor or biomarker has been definitively proven to be predictive. Instead, there is a growing interest in the gut microbiota to influence MTX bioavailability and as a predictor of subsequent clinical responses to MTX in the RA patients. The next section, we will detail the potential interaction between MTX and the gut microbiota.

3 PHARMACOMICROBIOLOGY OF MTX IN RA

Current treatment guidelines for RA [such as ACR (29) and EULAR (94)] recommend MTX for all patients with early RA, because early and aggressive MTX intervention can slow the onset and progression of RA, reduce disease activity, alleviate clinical symptoms, and even radiographic progression in some RA patients. Therefore, there is an urgent need for in-depth understanding to identify factors and predictors that influence MTX in order to maximize clinical efficacy and mitigate toxic effects, thereby eliminating frustration and reducing the waste of medical expenditures. Currently, a variety of candidate biomarkers have been considered to have potential predictive value for clinical efficacy of MTX, including clinical phenotypes, inflammatory pathway gene expression, host genetics, autoantibodies, and cytokines, etc, involving proteomics, metabolomics, transcriptomics and other omics. Unfortunately, none of them has been able to accurately predict clinical response to MTX (95).

A growing body of evidence suggests that non-human genetic factors, particularly from the trillions of microorganisms that is the microbiome, may contribute to the development of inflammatory arthritis in genetically susceptible individuals (96–98). Intriguingly, variability in interindividual microbiome composition and metabolic capacity play a unique role in determining the clinical efficacy (and the development associated with adverse events) of certain drugs (99–101). In turn, drugs may also exert their anti-inflammatory effects by altering a patient’s gut microbiome. The study of the interactions between drugs and microorganisms, known as pharmacobiology, has led us to the concept of precision medicine, which is not foreign to most rheumatologists. The long-term and ultimate goal of this discipline is to manipulate the complex host-associated microbiome (both in vivo and on the surface of the body), in order to predict clinical responses, improve clinical efficacy, and minimize adverse events (17).

3.1 Effect of Gut Microbiome on Clinical Efficacy of MTX in RA

What accounts for variation in clinical outcomes between individuals? The answer is not entirely clear. Moreover, to date, known factors such as host genetics, laboratory or host biomarkers cannot be used to accurately predict clinical response to a drug (95). Furthermore, differences in the metagenomes of microorganisms in the gut of RA patients such as bacterial strains, genes, enzymes, proteins and/or metabolites may at least partially determine their bioavailability and/or subsequent response to MTX. So, there is reason to believe that there is some connection between MTX and the gut microbiota.

3.1.1 Gut Microbiome Partly Determines How RA Patients Respond to MTX

A preliminary study using red blood cell MTX polyglutamate concentration explained 20% of the variation in MTX response, but unfortunately this finding was not consistently reproduced in other cohorts (102–105). Other possible factors including serum or plasma MTX concentration (106, 107), clinical factors such as sex and disease activity (92, 108–110) and circulating CD39+ regulatory T cells (92, 111), and genetic factors such as gene polymorphism (112) as predictors of MTX efficacy all failed. More than a decade ago, the first clinical pharmacological model (113) was proposed to predict the clinical response or not of MTX monotherapy to newly diagnosed RA. This model does not accurately predict the clinical efficacy of RA and cannot be widely used in the population, although it improves on the original genetic-based model by incorporating multiple variables (114–116).

The fact that so many other factors do not account for individual differences in MTX response raises the possibility that the differences are caused at least in part by individual differences in the metagenome of the gut microbiome. More recently, some research groups have suggested that intestinal microbiota is closely related to immunoregulatory therapy, and that it may have great predictive value in the clinical response of drugs (117–122). Microbial differences found in the gastrointestinal tract of RA patients may partially determine the bioavailability and/or subsequent clinical outcome of MTX. This theory was first demonstrated in rodents. Valerino, D. M., Johns, D. G. et al. (41) found that the intestinal absorption and metabolism of MTX in germ-free and antibiotic-treated mice were reduced compared to wild-type mice, suggesting that the biotransformation of MTX was hindered if the microbiome was not tasted, that is, gut microorganisms is an essential link in the absorption and metabolism of MTX. Moreover, according to a 2013 study (97) found that untreated new RA patients with gut microbiome differences between purine metabolic pathways, including four hydrogen folic acid (and other purines) biosynthesis, adjustable oral absorption and bioavailability of MTX and downstream treatment effect. Another study (123), which differentiated RA patients from healthy controls based on differences in oral microbiota, found that clinical measures such as disease activity and symptoms such as joint pain were strongly associated with changes in the microbiome. This suggests the potential diagnostic and prognostic value of the gut microbiota in RA. However, this study focused on the oral microbiome (which have a lesser similarity to gut microbiota).
as a predictor, focusing primarily on confirmed patients with a long-term RA, who had a significantly different microbiome from the new-onset (NORA) subjects.

A 2020 study (124) delved into the relationship between gut microbes and oral MTX response in NORA. They found that the gut microbial community structure before MTX administration, made the difference between clinical response or not, which is that MTX-responders (MTX-R) has significantly lower microbial diversity compared with non-responder (MTX-NR). At the phylum level, the ratio of Firmicutes/Bacteroidetes in MTX-NR is high. At the same time, MTX-rich bacteria in NR samples are: the Euryarchaeota phylum, unclassified Clostridiales/ Clostridiales Incertae Sedis XIII (family) and Escherichia/ Shigella. By contrast, in MTX-R, Prevotella and Bacteroides genera are significantly more abundant. The team then used shotgun sequencing to define differences in metagenome and gene abundance of gut bacteria between the two groups and found 6,356 KEGG Orthologs (KOs). 7 microbial modules (mainly involves metabolic and biosynthetic potential) and 462 KOs were identified for the separation of MTX-NR and MTX-R. For instance, many pathways including MAPK signaling pathway, fatty acid degradation and DNA replication were significantly increased in MTX-NR. The 2013 study (97) also distinguished MTX responders from non-responders in terms of microbial metabolic pathways. To sum up, we conclude gut bacteria metagenome including microbial or bacterial function may be more likely to be related to clinical responses than microbiol community structure.

Taken together, NORA patients who responded well to MTX had a different gut microbiome than MTX-NR, which allows us to predict the clinical absence of MTX by using pretreated microbiota.

3.1.2 Pharmacobiological Methods Can Be Used to Predict the Clinical Efficacy of MTX in RA Patients

Research groups studying human autoimmune diseases use pharmacobiological methods to analyze the gut microbiome and/or its gene coding function as predictors of biotherapy response (17). The use of gut microbiome to predict the clinical respond of MTX in newly diagnosed RA patients was included. These research groups came to two conclusions: gut metagenome at baseline could differentiate MTX responders from non-responders and another is that “ex vivo incubation with MTX of samples from patients with treatment-naive, new-onset RA correlated with the magnitude of future clinical response”. And accordingly, a small idea that a new and potentially valuable tool (metagenome-based classifiers) may be available for the decision-making of newly emerging RA was put forward. That’s exactly what the 2020 study has found.

Data from this 2020 study (124) demonstrate that gut metagenomics at baseline can predict MTX clinical response or not and provide a predictive model for it, as well as a possible plausible mechanism for the relationship between gut microbiome and clinical response to MTX prior to treatment. As described earlier, there were significant differences in gut metagenomes between MTX-NR and MTX-R patients prior to the start of treatment. Therefore, machine learning based on gut microbiome metagenomes can robustly predict MTX responses. The research group built a random forest model to predict whether MTX would respond or not. At the same time, they proved that a model based on microbiome characteristics, rather than clinical pharmacological characteristics, can determine response to MTX. However, the potential clinical application of this model is limited to drug-naïve and RA patients with direct exposure to MTX. Although this model has many deficiencies, it has contributed greatly to the prediction of clinical response of MTX by intestinal metagenomics at baseline compared to the previous model (113).

Intestinal metagenomes with significant differences before treatment initiation can predict clinical response to MTX. And why is this? Let’s put forward a hypothesis that the gut metagenomes are associated with known MTX metabolic pathways. And this hypothesis has been confirmed. As what mentioned before, a 2013 study (97) found that untreated new RA patients with gut microbiome differences between purine metabolic pathways. The MTX-NR microbiome showed the increased genes abundance was mainly concentrated in the genes encoding for purine nucleoside phosphorylase and adenosine deaminase, which increased hypoxanthine and reduced the bioavailability and cytotoxicity of MTX (125, 126). In contrast, the gene-coding enzymes (i.e. hisH and hisA) associated with the accumulation of 5-aminimidazole-4-carboxylate ribonucleoside (AICAR) were relatively reduced in MTX-NR (127). These two enzymes can regulate many immune-mediated effects of MTX by regulating the amount of AICAR. These two examples further highlight the importance of characterization of gut metagenomic, including enzymes or proteins, for the development of drug metabolic functions. Unfortunately, given that the gut metagenome contains a collection of all the genomes in the microbiome, it is difficult to determine the efficacy of MTX at the individual level and in clinical practice, although it can distinguish MTX significantly at the group level. If cost is taken into account, the actual situation becomes more complicated.

Then the group (124) used two independent metabolomic platforms to confirm the differences in MTX clinical responses were directly mediated by NORA gut microbiome via affecting its metabolism. This provides for the first time a possible mechanism for the association between gut microbiome and drug response, that is to say, “gut bacteria derived from MTX non-responders differentially deplete MTX ex vivo and remaining drug levels correlate with decreased clinical response”. First, the ability of the fecal microbiota to metabolize MTX in vivo is highly variable between individuals. Then, fecal microbiome in the MTX-NR group could rapidly reduce the concentration of MTX, while the concentration of MTX in the fecal supernatant of the MTX-R group remained unchanged. Last, the MTX elimination rate in each individual was significantly inversely correlated with future observed clinical responses. MTX levels measured in the supernatant related to future clinical responses when incubated with fecal samples. Therefore, patients’ gut microbiome may be a predictor of clinical response to MTX.
Taken together, we obtained the conclusion that gut microbiome (more accurately, gut microbiome metagenome) may serve as a predictor of clinical MTX response and the possible mechanism by which gut microbiome acts on MTX, where patients with more abundant gut bacteria were able to efficiently metabolize and/or consume MTX, and were associated with worsening clinical outcomes (see Figure 4).

3.2 Influence of Gut Microbiome or Its Metabolites on MTX Gastrointestinal Toxicity

Long-term oral administration of MTX may cause many adverse reactions, including intestinal toxicity, cardiac toxicity, bone marrow toxicity, renal toxicity and pulmonary toxicity, which greatly limits the clinical application of MTX (128, 129). Among them, intestinal toxicity is the most common side effect (18). The main symptoms include nausea, vomiting, abdominal pain, bloating, and diarrhea, leading to malabsorption, weight loss, and eventually discontinuation of chemotherapy (25, 128, 130). It is possible that MTX exerts toxic effects on normal intestinal cells by disrupting metabolic pathways such as anti-inflammatory, anti-oxidant and apoptosis (131). Another possible mechanism is that changes in gut microorganisms or their metabolites cause polarization of macrophages, leading to the release of inflammatory factors such as interleukin-12 (IL-12) and tumor necrosis factor (TNF), and ultimately aggravate gut mucosal injury. However, it is a complex process for MTX to cause gastrointestinal damage, which is not entirely clear so far. Therefore, there are no satisfactory therapeutic interventions to prevent or treat this adverse reaction.

The homeostasis of the intestinal environment is critical to the integrity and function of the entire gastrointestinal tract (132). The intestinal microbiota plays an important role in the development and progression of mucositis, which is the central manifestation of intestinal damage (133). Changes in the composition, structure and metabolites of the gut microbiome may affect the metabolism of MTX, thus affecting the status of gut mucosa. However, these complex dynamic processes need further investigation.

Some bacteria (18) or metabolites (134, 135) including amino acids, polyphenols or vitamins have been reported to influence MTX-induced intestinal mucosal injury. One possible explanation is that chronic exposure to MTX may induce community and functionality changes in the intestinal, and induce disruption of downstream CPDG2 activity, which may delay the detoxification of DAMPA by MTX, leading to increased gastrointestinal toxicity (19). However, each strain or metabolite administered individually has a limited range of effects on gastrointestinal toxicity. Therefore, it is feasible to reduce the gastrointestinal toxicity of MTX by establishing a combination therapy with candidate bacteria and metabolites (136). It has also been reported that changes in diet cause changes in the microenvironment of the gut microbiota, which ultimately aggravates MTX-induced intestinal inflammation (136). The proteins or lipids in high-fat high-sucrose diet (HFHSD) alter the intestinal environment and affect the pathogenesis of MTX-induced intestinal mucosal injury.

At present, treatment with leucovorin (LV) is the most common method to reduce the toxic effects of MTX. Previous studies (137, 138) have confirmed that the decrease in the severity of intestinal inflammation after the combined administration of MTX and LV is related to the diversity, richness and species composition of gut microbial community. But not all species of gut bacteria are affected to the same degree. A recent study (139) has shown that LV can improve MTX intestinal toxicity, possibly via increasing the composition of beneficial bacteria such as Bifidobacterium and reversing the imbalance of intestinal microbiota caused by MTX. A 2018 study...
experiment seems to disagree. This is a double-blind, parallel-
symptoms. The failure rate (i.e., patients requiring intensive
anaerobic environment (141), need to be further attended in
the control group. Therefore, other mechanisms, such as the
degree and persistence of donor microbiota implantation, and
the future.

Therefore, we speculate that
fecal microbiota transplantation (FMT) ameliorates drug-
reported that arti-
induced intestinal damage (140). Therefore, we speculate that
to reduce MTX-induced gut in-
probiotics and/or FMT may be a safer and more effective method
use metagenomic sequencing to
on MTX-induced in-
richness and diversity of the microbiota (123, 144), including
the humanized mice in dose dependent manner the composition
combined relative abundance (17).

MTX in both in vitro and the biological experiments can affect
the humanized mice in dose dependent manner the composition
of gut microbiome (19). Low doses of MTX increased the species
richness and diversity of the microbiota (123, 144), including
increasing relative abundance of Firmicutes and decreasing
relative abundance of Bacteroidetes, which reverse the
perturbations of the microbiota normally associated with RA
(17). However, high doses of MTX significantly reduced
intestinal bacterial diversity (145). Most of the changes were
due to a relative decrease in anaerobic bacteria of Firmicutes,
accompanied by a relative increase in Bacteroidetes. High doses
of MTX may alter the function or abundance of bacterial
enzymes involved in MTX hydrolysis, such as CPDG2, which
we mentioned earlier (145). Other studies have shown that MTX
exerts an anti-inflammatory effect via interacting with off-target
bacterial enzymes such as DHFR in E. coli or Lactobacillus Casey
(146). And low-dose MTX-induced changes in the gut
microbiome reduce host immune activation (147). Specifically,
the microflora changes lead to the reduction of multiple host cell
populations, such as activated T cells, B cells, myeloid cells,
etc. At the same time, reduced stimulation of immune system
activation and decreased tolerance to Treg cell induction
suggested that MTX-induced changes in the microbial
community reduced its possible inflammatory potential.

A recent study shows that gut microbiota is closely correlated
with P – glycoprotein (P-gp, a glycoprotein associated with MTX
efficacy) (148). For example, Lachnolosstrium is positively
correlated with P-gp, while Turicibacter is negatively correlated
with P-gp. As mentioned above, MTX has a significant impact on
the gut microbiome of RA patients according to its dose, and
combined with the relationship between gut microbiome and P-
gp, it is reasonable to speculate that oral MTX affects the
therapeutic effect of the drug itself by affecting the gut
microbiome. Unfortunately, so far, this idea has only existed in
theory and has not been confirmed experimentally.

The gut metagenome of RA patients before treatment
determines whether or not they respond to MTX, and that in
vitro incubation is correlated with the clinical response intensity.
In turn, different doses of MTX altered the species richness and
diversity of intestinal bacteria (see Figure 5). In other words,
there is a dose-dependent bidirectional interaction between MTX
and gut microbiota. This interaction between drugs and
microbes is known as pharmacobiology, a term not foreign
to rheumatologists.

3.3 Effects of MTX on Gut Microbiome in RA Patients

Gut microbiome can be used as a predictor of clinical response
to MTX, so, conversely, does MTX affect gut microbiome? The
answer is of course yes. According a using 16 s - seq study (142),
with the passage of time, the routine oral dose MTX will not lead
to a gut microbial ecology of persistent disturbances. But many
subsequent experiments have demonstrated that MTX can also
affect the community structure of intestinal microorganisms,
regulating their diversity and function (17). It has recently been
reported that MTX inhibits the growth of 30% of 40
representative intestinal bacterial strains and 84% of 43 bacterial
isolates (143), covering 43% of the human gut microbiota in their
combined relative abundance (17).

MTX in both in vitro and the biological experiments can affect
the humanized mice in dose dependent manner the composition
of gut microbiome (19). Low doses of MTX increased the species
richness and diversity of the microbiota (123, 144), including
increasing relative abundance of Firmicutes and decreasing
relative abundance of Bacteroidetes, which reverse the
perturbations of the microbiota normally associated with RA
(17). However, high doses of MTX significantly reduced
intestinal bacterial diversity (145). Most of the changes were
due to a relative decrease in anaerobic bacteria of Firmicutes,
accompanied by a relative increase in Bacteroidetes. High doses
of MTX may alter the function or abundance of bacterial
enzymes involved in MTX hydrolysis, such as CPDG2, which
we mentioned earlier (145). Other studies have shown that MTX
exerts an anti-inflammatory effect via interacting with off-target
bacterial enzymes such as DHFR in E. coli or Lactobacillus Casey
(146). And low-dose MTX-induced changes in the gut
microbiome reduce host immune activation (147). Specifically,
the microflora changes lead to the reduction of multiple host cell
populations, such as activated T cells, B cells, myeloid cells,
adjuvant probiotics and probiotics, which have shown positive results (137). New technologies such as organ-on-a-chip (150, 151) and bacterial culture (152) help to understand the mechanisms of pharmacomicrobiology via attempting to mimic the gut environment and reproduce the physiological interactions of host microbes. Then, the drugs to be studied are cultivated in these systems to explore the interaction between drugs and microorganisms, including the effects of drugs on bacterial metabolism and reproduction, and the effects and mechanisms of bacteria on drug biotransformation. These pharmacobiological studies will provide a new idea for the treatment of rheumatoid arthritis and even inflammatory arthritis.

4 CONCLUSIONS AND PERSPECTIVES

Although several efficacious therapies have recently been developed, MTX has always been the basic and first-line drug for the treatment of RA. Multiple mechanisms may contribute to the anti-inflammatory effects of MTX including inhibition of purine and pyrimidine synthesis, promotion of adenosine and some cytokines release, inhibition of ROS, etc., as well as transmethylation reaction, activator (STAT) pathway and expression of some long non-coding RNAs, which are not described in detail in this paper (3, 27). However, there is still a high proportion of RA patients who do not respond or have major toxic effects to MTX. We speculate that this inter-individual difference is at least partly related to the gut microbes of patients, and more and more trials have proved this speculation to be correct. In other words, gut microbes are associated with MTX resistance or poor response. Gut metagenome at baseline could differentiate MTX responders from non-responders and ex vivo incubation with MTX of samples from patients with treatment-naive, new-onset RA correlated with the magnitude of future clinical response. In turn, MTX affected the composition of intestinal microbes in humanized mice in a dose-dependent manner, inhibiting the growth of certain intestinal bacteria. Although comparison of MTX consumption rate and response probability by metagenome-based models did not find a significant correlation, this may indicate that most gene orthologs included in the model do not appear to play an important role in MTX depletion. A possible explanation is that these gene orthologs may affect the MTX response, but independently of drug metabolism. Future research needs to determine whether the microbiome can directly metabolize MTX in the body, jump-start the immune system to boost the response, or both. The rigorous integration of clinical factors, host genomics, and pharmacobiology will lay the foundation for major advances in rheumatic precision medical. More research is needed on the interactions between drugs and gut bacteria, known as pharmacobiology, which will be a research hotspot now and even in the future in the field of rheumatic diseases including RA.

AUTHOR CONTRIBUTIONS

HY drafted the manuscript, prepared illustrations and discussed the content with the other authors. CW conceived the topic and revised the manuscript. RS intellectually revised the manuscript. HX, CG and XL also critically revised the manuscript for intellectual content. All the authors approved the manuscript for publication.

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