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**optimal immunosuppressor induces stable gut microbiota after liver transplantation**

Jiang JW *et al*. Immunosuppressor and gut microbiota after transplantation

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**Abstract**

***AIM***

To study the influence of tacrolimus (FK506) with different dosages on gut microbiota after liver transplantation (LT) in rat.

***METHODS***

Specific pathogen-free Brown Norway (BN) rats and Lewis rats were used in this experiment with five groups: (1) Tolerance group (BN-BN LT, *n* = 8); (2) rejection group (Lewis-BN LT, *n* = 8); (3) high dosage FK506 (FK506-H) group (Lewis-BN LT, *n =* 8); (4) middle dosage FK506 (FK506-M) group (Lewis-BN LT, *n =* 8); (5) low dosage FK506 (FK506-L) group (Lewis-BN LT, *n =* 8). FK506 was administered to receipts at a dose of 1.0 mg/kg, 0.5 mg/kg, and 0.1 mg/kg weight for 30 d after LT to the FK506-H, FK506-M, and FK506-L groups, respectively. On the 30th day after LT, all rats were sampled and then euthanized. Blood samples were harvested for liver function and plasma endotoxin testing. Hepatic graft and ileocecal tissues were collected for histopathology observation. Ileocecal contents were used for DNA extraction, Real-time quantitative polymerase chain reaction (RT-PCR) and digital processing of denaturing gradient gel electrophoresis (DGGE) profiles and analysis.

***RESULTS***

Compared to the FK506-H and FK506-L groups, FK506-M was optimal for maintaining immunosuppression and induced normal graft function; the FK506-M maintained the integrity of gut barrier and low plasma endotoxin levels; furthermore, DGGE results showed that FK506-M induced stable gut microbiota; Diversity analysis indicated that FK506-M increased species richness and rare species abundance; Cluster analysis confirmed the stable gut microbiota induced by the FK506-M; Phylogenetic tree analysis identified crucial bacteria associated with the FK506-M, seven of the 9 bacteria that were decreased corresponded to *Bacteroidetes*, while increased bacteria were of the *Bifidobacterium* species; the FK506-M increased *Faecalibacterium prausnitzii* and *Bifidobacterium spp*. and decreased *Bacteroides-Prevotella* and *Enterobacteriaceae* as assessed with RT-PCR, which confirmed the crucial bacterial alterations identified through DGGE.

***CONCLUSION***

Compared to the low or the high dosage of FK506, an optimal dosage of FK506 induces immunosuppression, normal graft function and stable gut microbiota following LT in rats, the stable gut microbiota presented increased probiotics and decreased potential pathogenic endotoxin-producing bacteria. These findings provide a novel strategy based on gut microbiota for immunosuppressive dosage assessment for recipients following LT.

**Key words:** Liver transplantation; Graft function; Gut microbiota; Immunosuppressor; Tacrolimus; Rejection; Denaturing gradient gel electrophoresis

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**Core tip:** This study is the first time to illustrate the effects of Tacrolimus (FK506) with different dosages on gut microbiota following liver transplantation (LT) and indicates that an optimal dosage of FK506 induces effective immunosupression, good graft function and stable gut microbiota after LT in rat. Based on the relationship between gut microbiota and immunosuppressive dosage in this study, we can not only illustrate precise changes of gut microbiota given by FK-506 with different dosages following LT, but we also provide a novel monitoring strategy based on changes in gut microbiota for immunosuppressive dosage assessment in patients following LT.

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**INTRODUCTION**

Liver transplantation (LT) is considered the only definitive therapy for end-stage liver diseases, including liver cirrhosis, liver failure and hepatocellular carcinoma[[1](#_ENREF_1),[2](#_ENREF_2)]. Currently, the 1-year survival rate in patients following LT has reached 90% and the 5-year survival has reached 75%[[3](#_ENREF_3)], however, rejection after LT remains the major cause of hepatic graft dysfunction and is a life-threatening complication[4]. Treatment with inadequate immunosuppression might lead to a higher risk of rejection and even graft dysfunction, whereas excessive immunosuppression has been closely associated with a greater incidence of infection, sepsis, drug toxicity, cancer, chronic graft dysfunction, renal dysfunction, and even increased mortality[5-7], with bacterial infection being the most prevalent type of infection[8]. Thus, the optimal dosage of immunosuppressive medication is crucial for prevention of rejection, lessen IS toxicity and treatment of patients following LT[9].

Gut microbiota has been considered the most important micro-ecosystem that possesses a symbiotic relationship within the body[10-12]. The gut microbiota plays a crucial role in the development of fatty liver disease[13], liver cirrhosis[[14](#_ENREF_13)], hepatocellular carcinoma[[15](#_ENREF_14)], liver ischemic injury[[16](#_ENREF_15)] and liver graft rejection[[17](#_ENREF_16)]. Our previous study indicated that the healing of hepatic injury could ameliorate gut barrier function and promote gut microbial restoration following LT in rat[[16](#_ENREF_15)]. We also found that graft rejection after LT could induce gut microbial alterations, thereby inducing subsequent gut barrier dysfunction, gut barrier dysfunction presented a decrease of fecal secretory immunoglobulin A (sIgA) and increase of blood endotoxin and tumor necrosis factor-α, RT-PCR results showed that genus *Faecalibacterium prausnitzii* and Lactobacillus were decreased, whereas Clostridium bolteae was increased, which might in turn aggravate hepatic rejection[18].

Nevertheless, the influence of immunosuppressive medication on gut microbiota following LT remains unclear, the association between the dosage of IS and gut microbial alterations requires urgent elucidation. In this study, we studied the influence of FK506 with different dosages on hepatic graft function and gut microbiota following LT in rat. Furthermore, we identified the gut microbial profile and crucial bacterial community constituents using DGGE, and further verified the alterations of dominant gut bacterial populations with RT-PCR.

**MATERIALS AND METHODS**

***Animals***

Specific pathogen-free (SPF) male inbred Brown Norway (BN) and Lewis rats (weight 220-250 g, age 12-15 wk) were purchased from Beijing Vital River Laboratories (Beijing, China)，they were housed in a clean-level animal house located at the First Affiliated Hospital, School of Medicine, Zhejiang University. All rats were housed at 22-24 °C in 12 h light/dark cycles and fed sterilized standard rat chow and water.

***Experimental design and protocol***

The study animals were divided into 5 groups as follows: (1) Tolerance group (BN-BN LT, *n =* 8); (2) rejection group (Lewis-BN LT, *n =* 8); (3) high dosage FK506 (FK506-H) group (Lewis-BN LT, *n =* 8); (4) middle dosage FK506 (FK506-M) group (Lewis-BN LT, *n =* 8); (5) low dosage FK506 (FK506-L) group (Lewis-BN LT, *n =* 8). In Tolerance group, both donors and recipients were BN rats; in the other four groups，both donors were Lewis and recipients were BN rats, FK506 was administered to receipts at a dose of 1.0, 0.5, and 0.1 mg/kg weight for 1 mo after LT to the FK506-H, FK506-M, and FK506-L groups, respectively. To mimic the clinical application, FK506 was administrated *via* abdominal subcutaneous injection, once every 12 h for 7 d after LT, and then *via* intragastric administration, once per day, for the following 8-30 d. The sustained-release FK506 were administered intragastrically to maintain a consistent dosage effect at 24 h.

The study was conducted in accordance with the ‘‘Guide for the Care and Use of Laboratory Animals’’ published by the National Institutes of Health (NIH publication 86-23, revised 1985). The experimental protocol was approved by the Animal Care and Use Committee of the First Affiliated Hospital, School of Medicine, Zhejiang University.

***Liver transplant surgical procedures and sample collection***

The LT surgery was performed according to our previous methods[[16](#_ENREF_15),19], with slight modifications. The anesthesia was performed by intraperitoneal injection of Ketamine Hydrochloride (100 mg/kg) and Atropine (1 mg/kg) (Shanghai No. 1 Biochemical and Pharmaceutical, China), and then ether was inhaled to maintain anesthesia[[6](#_ENREF_15)]. All recipients were revived shortly after the procedure and no further treatment was administered.

All rats were sampled at 30th day after LT. The abdominal aorta was punctured and blood samples were harvested for liver function and plasma endotoxin testing. Hepatic graft and ileum tissues near ileocecus were collected for morphological observation. Ileocecal contents were harvested and stored at -80 °C for gut microbial analysis. All rats were then euthanized *via* an overdose of anesthetic.

***Liver function and*** ***plasma endotoxin testing***

Plasma alanine aminotransferase (ALT) and aspartate aminotransferase (AST) levels were detected using an automatic biochemical analyzer (Hitachi 7600, Tokyo, Japan).Plasma endotoxin level was determined using a colorimetric Limulus Test (Shanghai Yihua Medical Technology Co., Ltd, China) in accordance with the manufacturer's instructions.

***Histopathology and Transmission electron microscopy (TEM) evaluation***

The hepatic graft and an ileum tissue sample taken 3 cm from the ileocecus were fixed in 40 g/L neutral formaldehyde and embedded in paraffin, cut into 4-μm slices, stained with hematoxylin and eosin (HE), and then analyzed under light microscopy**.** At the same time, the same samples about 1mm3 were fixed in a 2.5% glutaraldehyde solution and prepared in accordance with the standard technical procedures for TEM, as previously described[20]. Hepatic and ileal ultrastructures were analyzedat the Imaging Facility of Core Facilities, Zhejiang University School of Medicine. Histopathology and TEM evaluation were investigated by a pathologist blinded to the treatment group.

***DNA extraction of ileocecal content***

DNA extraction of a frozen aliquot of each ileocecal fecal sample was performed as in our previous studies[[16](#_ENREF_15)]. DNA concentration was measured by NanoDrop (Thermo Scientific) and DNA integrity was verified using agarose gel electrophoresis.

***Real-time quantitative PCR***

The primers used for quantitative RT-PCR (RT-qPCR) are listed in Supplementary Table 1. All oligonucleotide primers were synthesized by TAKARA (Dalian, China). RT-qPCR was performed as reported previously[[21](#_ENREF_20)]. The copy number of 16s rDNA operons per microliter of crude DNA template was determined by comparing serially diluted plasmid DNA standards running on the same plate.

***DGGE profiling***

The V3 variable region of 16S rDNA was amplified using the hot-start touchdown protocol with specific primers for the conserved regions of 16S rDNA. DGGE was performed using the D-Code universal mutation detection system apparatus (Bio-Rad, Hercules, CA) with 16 cm × 18 cm × 1.5 mm gels according to the manufacturer's protocol. On each DGGE gel, standard references in the middle and two at both ends were used for digital gel normalization and comparison of gels.

***Digital processing of DGGE profiles***

DGGE profiles were digitally processed using BioNumerics software version 6.01 (Applied Maths, St-Martens-Latem, Belgium) in a multistep procedure following the manufacturer’s instructions. The parameters for band-class allocation followed those of our previous study[[16](#_ENREF_15)]. Quantitative information of a given band was calculated using the Gel-Pro analyzer 4 software (Media Cybernetics, USA). Diversity was calculated using Shannon’s diversity, Simpson index, species richness, Chao-1, Fisher alpha, and Menhinick index with Past software (<http://folk.uio.no/ohammer/past/>)[[21](#_ENREF_21)]. Cluster analysis of the DGGE was conducted with an unweighted pair-group method with arithmetic means (UPGMA) based on the Dice similarity coefficient (band-based). Multidimensional scaling (MDS) and principal components analysis (PCA) were utilized following the instructions of the BioNumerics software.

***Sequencing of DGGE bands***

Specific DGGE bands of interest were excised and sequenced. Positive clones were verified and sequenced using Sanger’s method on an ABI 3730 automated sequencing system (Invitrogen, Shanghai, China). Homology searches of the GenBank DNA database were performed using the BLAST tool. Reference sequences of phylogenetic neighbor species (up to 97% similarity) were included to construct a phylogenetic tree with MEGA 5.0 program based on the neighbor-joining method[[15](#_ENREF_15)].

***Statistical Analysis***

Continuous variables were summarized as mean ± standard deviation (SD). One-way variance analysis followed by Dunn’s multiple comparison tests was utilized to evaluate statistical differences among different groups. Student *t*-test was used to examine parametric data between the two groups. Statistical analyses were performed using SPSS version 19.0 for Windows (SPSS Inc., Chicago, IL, United States). A *P*-value of less than 0.05 was considered statistically significant.

**RESULTS**

***A middle dosage of FK506 is optimal for maintaining liver transplant immunosupression***

An overdose or insufficient dosage of FK506 may lead to graft dysfunction, thus we first observed survival of all recipients. All rats following LT survived except one LT recipient in the FK506-H group that died of severe abdomen infection, which suggested an over-suppression of the immune status, thereby inducing infection and increasing mortality.

To illustrate the influence of different dosages of FK506 on hepatic grafts, we observed graft morphology. Under light microscopy (Figure 1A), hepatic grafts in the tolerance group presented a regular structure with well-arranged hepatocyte cords. In the rejection group, graft histology showed no obvious hepatocyte cords, accumulation of numerous red blood cells, and marked hepatocyte necrosis. However, FK506-M treatment preserved an approximately normal hepatic structure without hepatic acute rejection. Notably, hepatic graft resulted in hepatocyte cord disruption, widened sinusoids, and middle rejection injury in the FK506-L dosage group, suggesting an insufficient suppression of the immune status, thereby inducing rejection. Thus, compared to the FK506-H and FK506-L groups, FK506-M was the optimal dosage for maintaining immunosupression in rats following LT.

We next observed hepatic graft ultrastructure with TEM (Figure 1B). In the tolerance group, a normal hepatocyte structure with intact mitochondria and endoplasmic reticulum was observed. Hepatic graft presented significant karyopyknosis, marked mitochondrial vacuolar degeneration, and organelle breakdown in the rejection group. Nevertheless, FK506-M treatment noticeably improved hepatocyte structure, mitochondria morphology, and other organelle structure. Notably, hepatocyte apoptosis and mild organelle damage were noted in the FK506-L group. Plasma ALT and AST levels reflected graft function. Compared with the rejection group, plasma ALT and AST were significantly decreased in the FK506-M treatment group (both *P* < 0.001). In addition, plasma ALT and AST were reduced in the FK506-M group versus the FK506-L group (both *P* < 0.05). There was no statistically significant difference between the tolerance and FK506-M-treated groups (Figure 1C).

***An optimal dosage of FK506 kept the gut barrier intact and maintained low endotoxin levels following LT***

The liver-gut circulation and the gut-liver axis are closely associated with gut and liver function following LT. Thus, we next observed the ileal mucosa ultrastructure using TEM (Figure 2A). In the tolerance group, intestinal epithelial cells presented a normal morphology with many homogenously distributed microvilli and integrated tight junctions. In contrast, microvilli loss, tight junction damage, and bacterial invasion in epithelial cells were observed in the rejection group. However, FK506-M treatment significantly improved intestinal epithelial structure and kept microvilli and tight junctions intact. Notably, epithelial cell damage, microvilli disruption, and wider lateral spaces between neighboring cells were observed in the FK506-L treatment group.

Plasma endotoxin reflects gut barrier function. Compared with the tolerance and FK506-M groups, plasma endotoxin was significantly increased in the FK506-H group (*P* < 0.01 and 0.05, respectively), suggesting that over-suppression of immune status might lead to higher endotoxin levels. Meanwhile, plasma endotoxins were remarkably elevated in the rejection and FK506-L groups compared to the tolerance group (both *P* < 0.001). However, FK506-M administration significantly decreased endotoxin levels compared to the rejection and FK506-L groups (both *P* < 0.001) (Figure 2B).

***An optimal dosage of FK506 induced a stable gut microbiota as determined by DGGE***

Endotoxins are the product of common gram-negative bacteria and can initiate various pathophysiological cascades[[22](#_ENREF_22)]. Plasma endotoxin changes are mainly derived from changes in the gut microbiota. Based on the DGGE method, we analyzed the distribution of the ileocecal bacterial community (Figure 3). To characterize the DGGE profiles, we utilized the Dice coefficient and UPGMA as a cluster method to indicate band similarity. These profiles formed two primary clusters. The left cluster mainly consisted of samples from the tolerance group, in which total similarity was 80.1%, with the similarity among lanes ranging from 80.1% to 91.1%. The other cluster contained samples mainly from the rejection group and the different doses of the FK506 groups. Importantly, the similarity among lanes ranged from 82.4% to 96.7% in the rejection group and from 82.7% to 95.4% in the FK506-M treatment group, suggesting a significant uniqueness and stability of the gut microbiota.

In contrast, the DGGE lanes from the FK506-H and FK506-L treatment groups presented an inordinate distribution and an evident difference. The similarity of these lanes was relatively low and could not be clustered together. These results based on the DGGE profile indicated that an optimal dosage of FK506 induced a unique and stable gut microbiota in rats following LT.

***Diversity analysis and cluster analysis of gut microbiota using DGGE profiles***

Gut microbial diversity and species richness are important indexes used to evaluate gut microbial balance and stability. Thus, we calculated the gray amount of each band in each lane of the DGGE profiles with Gel-Pro analyzer to compare microbial diversity. There were no statistical differences in the Shannon and Simpson indexes among the five groups (Figure 4A). Compared to the tolerance group, species richness was significantly decreased in the FK506-H, FK506-L, and rejection groups (all *P* < 0.05). In contrast, FK506-M treatment markedly increased species richness compared to the FK506-H, FK506-L, and rejection groups (*P* < 0.01, 0.05 and 0.001, respectively) (Figure 4A).

We also analyzed the abundance of rare species, estimated using the Chao-1, Fisher alpha, and Menhinick indexes. Compared with the tolerance group, Chao-1, Fisher alpha and Menhinick indexes were remarkably reduced in the FK506-H, FK506-L, and rejection groups (all *P* < 0.05). However, FK506-M treatment significantly increased the abundance of rare species in contrast to the FK506-H, FK506-L, and rejection groups (all *P* < 0.05) (Figure 4B).

Furthermore, based on DGGE profiles, we conducted MDS and PCA analysis. The Euclidean distance between two points indicates the similarity between the two. Gut microbial structures from samples of the tolerance, FK506-M, and rejection groups were respectively clustered together, and showed separation from each other using MDS analysis (Dim 1, Dim 2, and Dim 3) (Figure 4C), suggesting a relatively unique and stable gut microbiota. PCA is an alternative approach to visualizing relationships among lanes using lane data (band classes). Our PCA analysis also gave similar results based on PCA axes X/Y/Z (13.7%, 11.2%, and 10.0%, respectively) (Figure 4D).

***Phylogenetic tree analysis of sequences based on DGGE profiles***

Of the 49 PCR-DGGE bands analyzed in the study, 48 band classes were selected. To identify crucial bacterial populations induced by FK506-M treatment, we compared band intensity between the FK506-M and the rejection groups. Of the 41 band classes selected, 28 showed slight variation in band intensity between the 2 groups. In contrast to the rejection group, the intensities of 9 band classes [18.2 (band 1), 19.9 (band 2), 61.0 (band 9), 72.9 (band 15), 81.0 (band 16), 82.0 (band 26), 85.3 (band 17), 86.5 (band 18), and 88.2 (band 19)] were significantly decreased (Figure 5A), while 4 band classes [33.7 (band 20), 41.8 (band 39), 56.8 (band 41), and 82.7 (band 47)] were increased in the FK506-M treated group (Figure 5B).

To determine the bacterial phylogenetic relationships and identify key bacteria involved in the microbial changes induced by FK506-M, the phylogenetic tree of sequences derived from DGGE bands was analyzed using the MEGA 5 software (Figure 6**)**. Almost all the matched bacteria of the 48 band classes were assigned to 3 phyla: *Bacteroidetes* (50.0%), *Firmicutes* (39.6%), and *Gammaproteobacteria* (10.4%). The closest matched bacteria shown on the phylogenetic tree corresponded to 13 crucial band classes associated with FK506-M treatment. Details of the 13 crucial band classes are presented in Supplementary Table 2. In these crucial bacteria associated with FK506-M treatment, 7 of the 9 decreased band classes were matched to *Bacteroidetes bacterium*, belonging to phylum *Bacteroidetes*, but the increased band classes were matched to the *Bifidobacterium species.*

***Quantitative verification of the predominant bacterial community in fecal microbiota using RT-qPCR***

To verify crucial bacteria alterations induced by FK506-M treatment, we further analyzed dominant bacterial populations using RT-qPCR (Figure 7). Compared with the tolerance group, *Faecalibacterium prausnitzii* and *Bifidobacterium spp.* were significantly decreased, whereas the *Bacteroides-Prevotella* group and *Enterobacteriaceae* were increased in the FK506-L treated and rejection groups (all *P* < 0.05). However, FK506-M treatment significantly increased *Faecalibacterium prausnitzii* and *Bifidobacterium spp.* and decreased the *Bacteroides-Prevotella* group and *Enterobacteriaceae* when compared to the FK506-L and rejection groups (all *P* < 0.05) (Figure 7A). These results essentially verified the changes observed in the levels of 13 crucial bacteria associated with FK506-M treatment based on the DGGE profiles.

Moreover, in the rejection group, the *Clostridium clusters I* was elevated, whereas the *Clostridium clusters XI* was reduced, in contrast to the other groups (Figure 7B). *Enterococcus faecalis* was specifically decreased in tolerance group (all *P* < 0.001). There were no statistical differences in the *Clostridium cluster XIVab* and *Lactobacillus* between the different groups (Figure 7C).

**DISCUSSION**

Liver transplantation is a life-saving technique for patients with end-stage liver disease. Appropriate dosage of immunosuppressive medications including FK506 could effectively prevent and treat allograft rejection in patients following LT. FK506 is a calcineurin inhibitor, which has become the most used immunosuppressant following LT[23]. Compared to cyclosporine, FK506 is more effective in reducing rejection, leading to better graft function and overall survival in LT patients[24]. Insufficient dosage of FK506 may lead to rejection and even graft dysfunction, while over dosage is closely associated with infection, increased complications，and even mortality, unfortunately, FK506 related toxicity, mainly due to overload usage is frequent in liver transplant patients[25].

In patients following LT, an optimal immunosuppressive therapy should balance the risk of infection, cancer, and drug toxicity caused by excessive immunosuppression, and the risk of rejection caused by an inadequately suppressed immune system[5]. To explore the optimal immunosuppressive agents dosage is very important for the long-term survival of the recipient and the liver graft. Jia[9] proved that compared to high FK506 blood concentration(10-15ng/L) with adverse effects such as infection, early renal impairment, hepatocellular carcinoma recurrence，even death, reasonable low FK506 blood concentration(5-10ng/ml) by less oral dosage especially in the early phase, namely “Minimizing tacrolimus”，could prevent graft rejection, avoid related toxicity, protect the graft function, and promote long-term survival of the recipient. Moreover，Geng[26] proved that lower FK506 trough levels (< 5ng/mL) in the late period of living donor liver transplantation are safe, improve the long-term outcomes.

Our previous studies indicated that bacterial community alterations in the gut were associated with liver cirrhosis[[14](#_ENREF_13)], hepatic graft rejection[18], hepatic injury[19], and pancreatic carcinoma[[27](#_ENREF_23)], however, the influence of immunosuppressive medication on the gut microbiota and the association between the dosage of immunosuppressive drugs and alterations in gut microbiota have not been reported to date. In our study, all recipients survived following LT, except one recipient in the FK506-H dosage group that died of severe abdomen infection, suggesting an over-suppression of the immune status. Hepatic graft presented intermediate rejection injury in the FK506-L group, suggesting insufficient immunosuppression. In contrast, the FK506-M dosage maintained approximately normal hepatic structure and function. Survival results and graft morphology indicated that FK506-M treatment was optimal for maintaining immunosuppression. Meanwhile, the FK506-M dosage kept the gut barrier intact and maintained low levels of plasma endotoxin. Thus, the FK506-M dosage was considered optimal for maintaining immunosupression in rats following LT.

Furthermore, we observed the influence of FK506 with different dosages on the gut microbiota. We further analyzed gut microbiota using a DGGE approach and found that the FK506-M dosage induced a unique and stable gut microbiota. MDS and PCA analysis validated the stable gut microbiota identified by DGGE. Phylogenetic tree analysis identified crucial bacteria associated with the FK506-M treatment. In addition, treatment with the FK506-M significantly increased the growth of *Faecalibacterium prausnitzii* and *Bifidobacterium spp*. and decreased that of the *Bacteroides-Prevotella* group and *Enterobacteriaceae* as determined *via* RT-qPCR, which essentially verified the alterations in the population of crucial bacteria observed from the DGGE profiles.

Endotoxin, also known as lipopolysaccharide (LPS), is the main product of common gram-negative bacteria and is a crucial factor for the association of gut microbiota with liver inflammation[[22](#_ENREF_22)]. In a pathogen-associated molecular pattern of microbiota-liver axis, LPS may possess the capacity to activate inflammation and initiate various pathophysiological cascades. High levels of LPS activate the NF-κB pathway, produce proinflammatory cytokines [tumor necrosis factor (TNF)-α, interleukin (IL)-6, and IL-1], and lead to liver inflammation and aggravation of hepatic oxidative damage[[28](#_ENREF_30)]. In our study, plasma endotoxins were remarkably elevated in the rejection and FK506-L dosage groups, but the FK506-M dosage significantly decreased endotoxin levels. To explore the causes of the observed endotoxin changes, we analyzed gut microbial alterations and identified the crucial bacteria involved. Importantly, compared to the rejection and FK506-L dosage groups, RT-qPCR results verified that FK506-M treatment significantly increased *Faecalibacterium prausnitzii* and *Bifidobacterium spp*. and decreased the growth of the *Bacteroides-Prevotella* group and Enterobacteriaceae, which are the main common gram-negative bacteria producing LPS. Thus, we speculated that the optimal dosage of FK506 might lead to an increase in probiotics and a decrease in the potential pathogenic endotoxins-producing bacteria.

In patients following LT, blood concentrations of immunosuppressant and its stability are important. To mimic clinical application, FK506 was administered *via* abdominal subcutaneous injection for 7 d and then *via* intragastric administration during the subsequent 8-28 d after LT. In patients, routine monitoring of blood tacrolimus concentration (TC) was performed using the PRO-TracTMⅡTacrolimus Elisa Kit (Diasorin, United States), however, there were no kit used for rat blood TC testing. According to our experimental experience, it may be due to immunological reasons, the blood TC in rats can not be tested by the above kit used for human. Fortunately, according to clinical practice of LT, different dosages of FK506 are positively correlated to different serum concentrations across a specific dosage range, meanwhile, intragastric dosages of FK506 were more feasible and easier to control the blood TC. In this experiment, the over-low dosage of FK506 led to a higher risk of rejection, whereas over-high dosage was closely associated with a greater incidence of infection, sepsis, and even increased mortality. Therefore, treatment with different dosages of FK506 instead of relying on the examination of blood TC was feasible and credible in rats after LT.

Gut microbiota alterations are closely associated with liver function in health and disease due to liver-gut circulation and the gut microbiota-liver axis[[29](#_ENREF_24)]. Gut microbiota is involved in a variety of human liver diseases, such as alcoholic liver disease[30,31], non-alcoholic fatty liver disease[32], non-alcoholic steatohepatitis[33], hepatitis B infection[[21](#_ENREF_20)], liver cirrhosis[[14](#_ENREF_13),34], and hepatocellular carcinoma[[15](#_ENREF_14),[28](#_ENREF_30),35]. Liver diseases always reflect changes due to alterations in intestinal permeability and gut microbial composition[[29](#_ENREF_24)], while an imbalance in the gut microbiota in turn, has been reported to be an important contributor to the progression of liver disease or liver injury[[36](#_ENREF_32)]. Notably, the improvement of hepatic injury might ameliorate gut barrier function and promote gut microbial restorations following LT, which might further benefit hepatic graft by positive feedback of the “gut-liver axis”[[16](#_ENREF_15),18]. Thus, the gut microbiota becomes an additional therapeutic target to improve hepatic injury after LT. In the present study, the optimal dosage of FK506 led to good hepatic graft function, kept the gut barrier intact, and induced a unique and stable gut microbiota, which further verify the effects of the “gut microbiota-liver” axis.

In conclusion, an optimal dosage of FK506 induces effective immunosupression, normal graft function and stable gut microbiota after LT in rat. A stable gut microbiota resulted in increased probiotics, including *Faecalibacterium prausnitzii* and *Bifidobacterium spp*. and decreased potential pathogenic endotoxin-producing bacteria, such as the *Bacteroides-Prevotella* group and Enterobacteriaceae. These findings provide a novel strategy involving the use of the gut microbiota for the assessment of the dosage of immunosuppressive medications and its effects in recipients following LT.

**ARTICLE HIGHLIGHTS**

***Research background***

Liver transplantation (LT) is the only definitive therapy for end-stage liver diseases. The optimal dosage of immunosuppressive medication is crucial for prevention of rejection, lessen side effects of immunosuppressor (IS) and treatment of patients following LT. The gut microbiota plays a crucial role in the development of obesity, diabetes, liver diseases and so on. Nevertheless, the influence of IS on gut microbiota following LT remains unclear, the association between the dosage of IS and gut microbial alterations requires urgent elucidation.

***Research motivation***

Acute rejection is still a leading cause of hepatic graft dysfunction following LT. Each recipients of LT must take IS to prevent acute rejection, but the effect of IS on gut microbiota remains unclear. Tacrolimus (FK506) is the main IS following LT, so we studied the influence of FK506 with different dosages on gut microbiota after LT in rat.

***Research objectives***

In this study, we studied the influence of FK506 with different dosages on hepatic graft function and gut microbiota following LT in rat. Furthermore, we will identify the precise changes of the gut microbiota given by different doses of FK-506 and provide new monitoring strategy for IS dosage assessment in recipients after LT.

***Research methods***

We underwent LT experiments in rat by taking different dosage of FK506 for 30 d, on the 30th day after LT, all rats were sampled and then euthanized.

We studied the hepatic graft function using serum ALT and AST examination and morphology change using histopathology and transmission electron microscopy evaluation. We also identified the gut microbial profile and crucial bacterial community constituents using digital processing of denaturing gradient gel electrophoresis (DGGE), and further verified the alterations of dominant gut bacterial populations with RT-PCR.

***Research results***

Compared to the FK506-H and FK506-L groups, FK506-M was optimal for maintaining immunosuppression and induced normal graft function; the FK506-M maintained the integrity of gut barrier and low plasma endotoxin levels; furthermore, FK506-M induced stable gut microbiota, increased species richness and rare species abundance by DGGE and Cluster analysis; The FK506-M increased *Faecalibacterium prausnitzii* and *Bifidobacterium spp*. and decreased *Bacteroides-Prevotella* and *Enterobacteriaceae* by Phylogenetic tree analysis and RT-PCR verification.

***Research conclusions***

An optimal dosage of FK506 induces effective immunosupression, normal graft function and stable gut microbiota after LT in rat. A stable gut microbiota resulted in increased probiotics and decreased potential pathogenic endotoxin-producing bacteria. These findings provide a novel strategy involving the use of the gut microbiota for the assessment of the dosage of immunosuppressive medications and its effects in recipients following LT.

***Research perspectives***

This study is the first to illustrate the effects of FK506 with different dosages on gut microbiota following LT and indicates that an optimal dosage of FK506 induces effective immunosupression, good graft function and stable gut microbiota after LT in rat. Based on the relationship between gut microbiota and immunosuppressive dosage in this study, we can not only illustrate precise changes of gut microbiota given by FK-506 with different dosages following LT, but we also provide a novel monitoring strategy based on changes in gut microbiota for IS assessment in recipients following LT. With the improvement of metagenomics and metabolomics techniques, the integrative study can further reveal that how gut microbiota participates in the side effects of IS on recipients, and gut microbiota may be a target to reduce side effects of IS. In addition, in order to further study the mechanism of the involvement of gut microbiota on the side effects of IS on recipients, it is essential to do clinical research.

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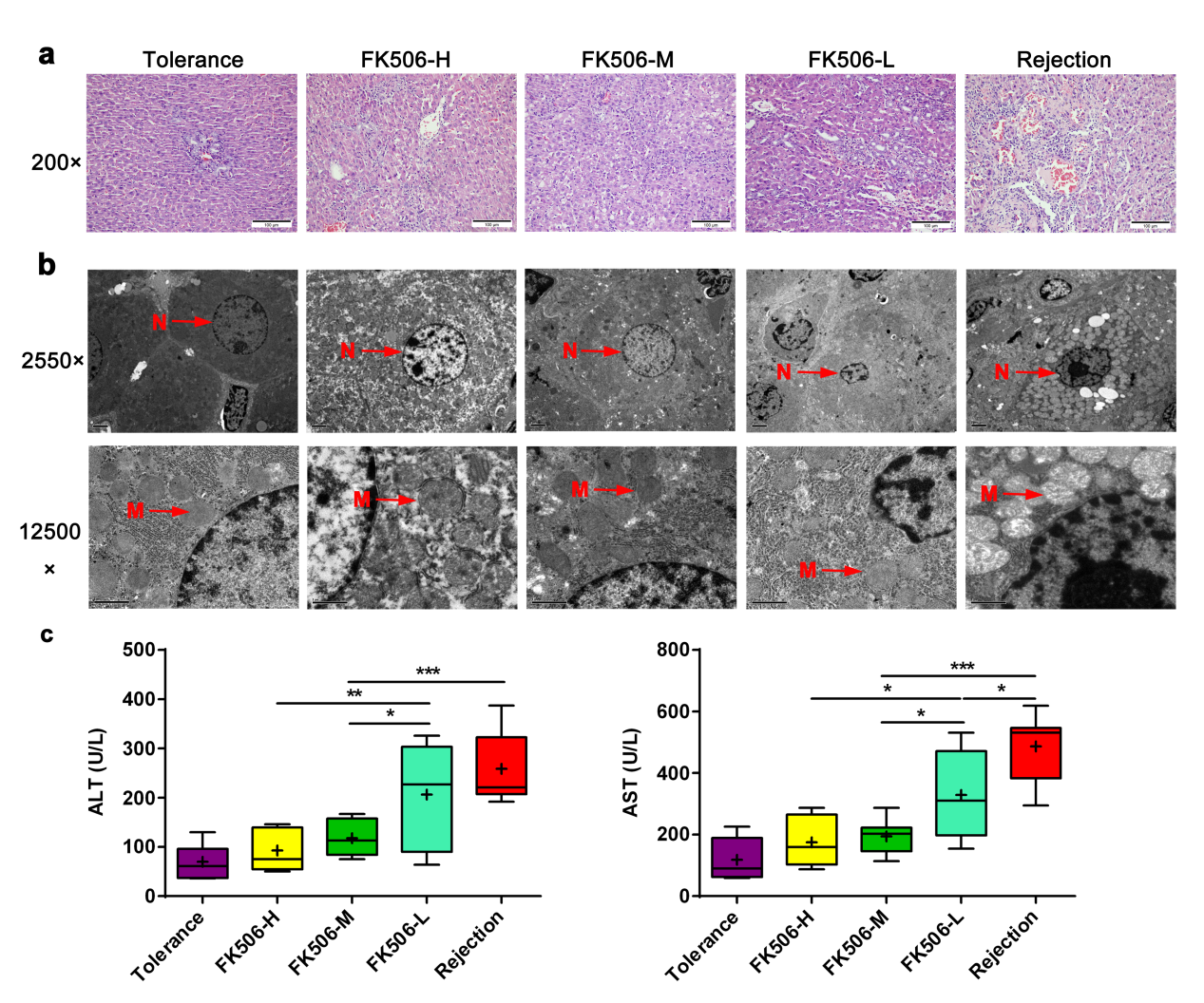
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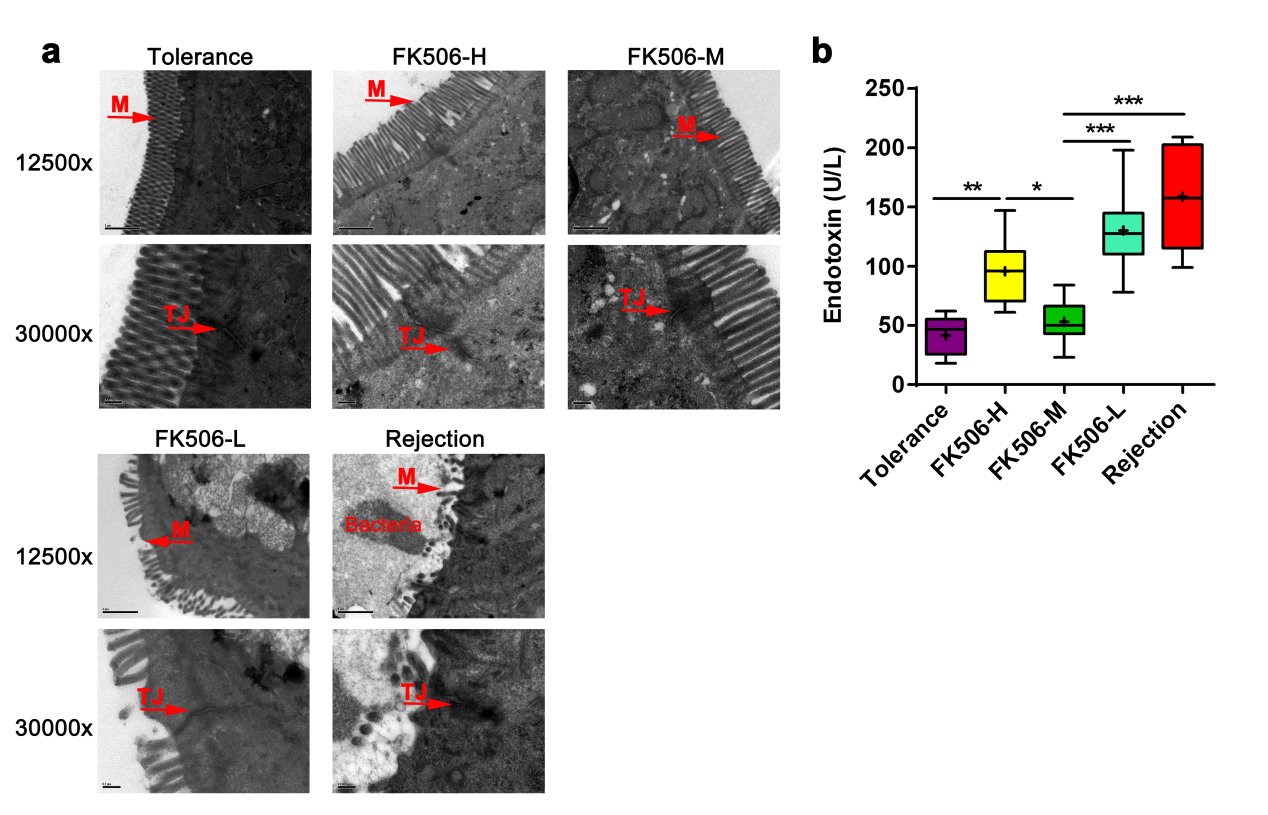
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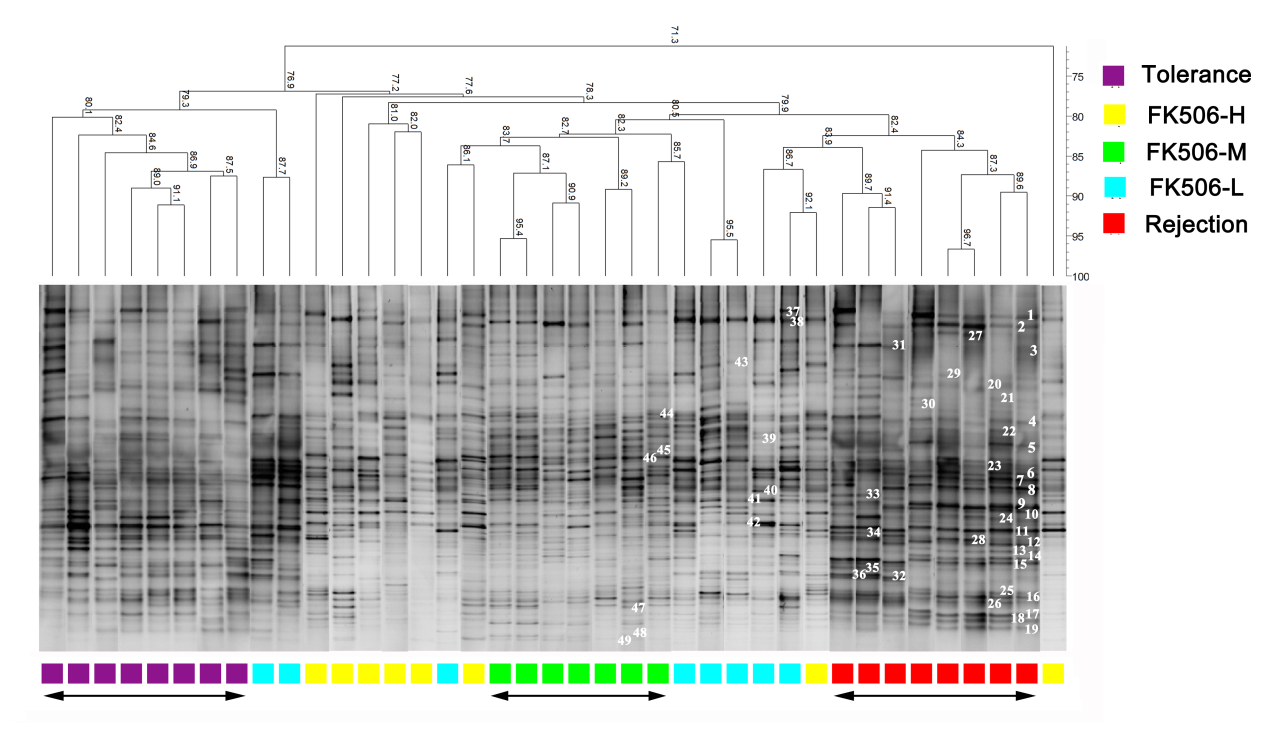
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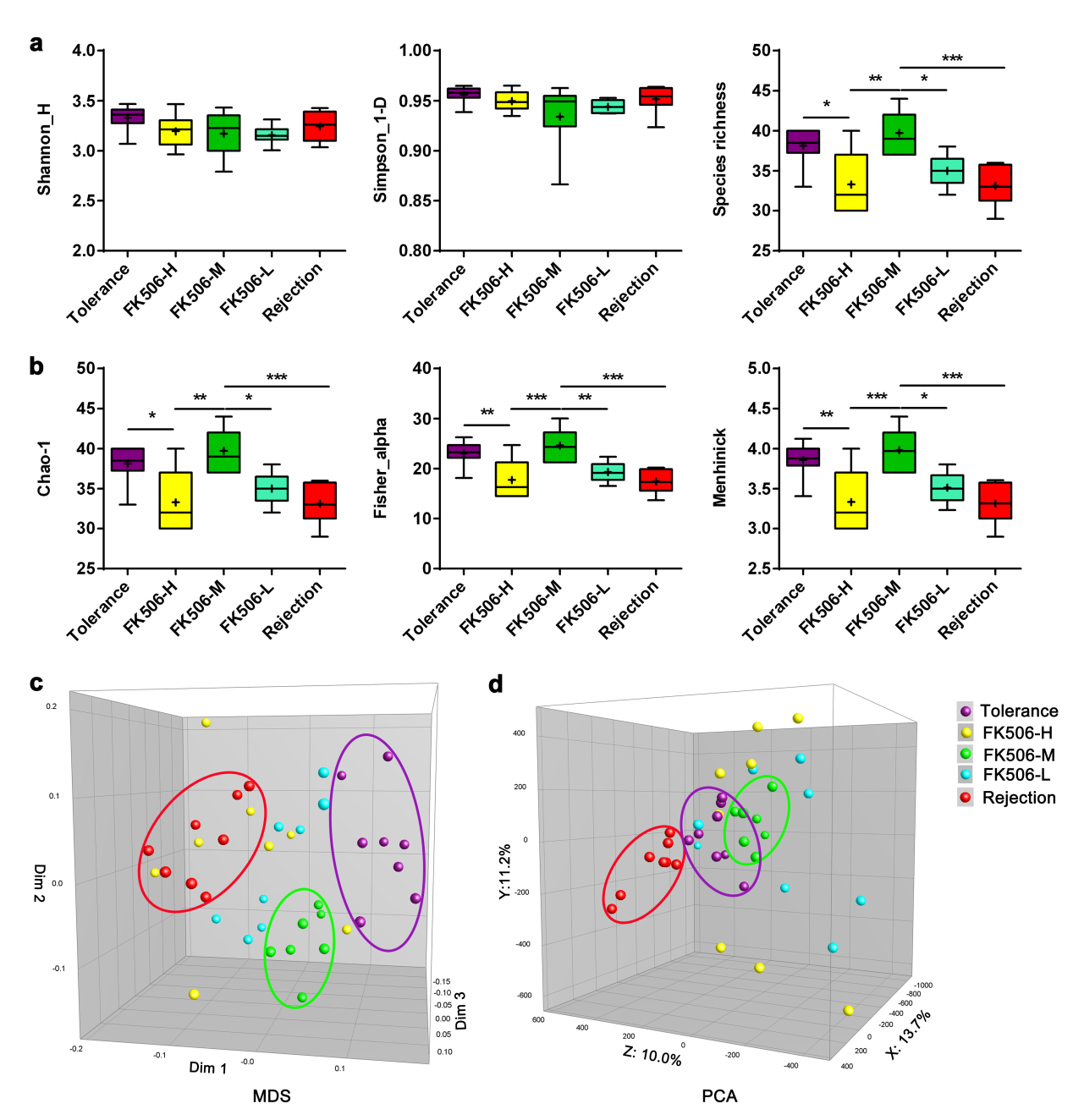
**Figure 1 Middle dosage of FK506 is optimal for maintaining immunosupression after liver transplantation. A:** Representative image of the hepatic graft pathological structure stained with hematoxylin and eosin (HE, 200 ×); B: Representative hepatocytes ultrastructure obtained *via* transmission electron microscopy (TEM, 2550 ×, 12500 ×); C: Assessment of hepatic graft function by plasma levels of alanine aminotransferase (ALT) and aspartate aminotransferase (AST) in different treatment groups. N: Cell nucleus; M: Mitochondria. a*P* < 0.05; b*P* < 0.01; c*P* < 0.001.

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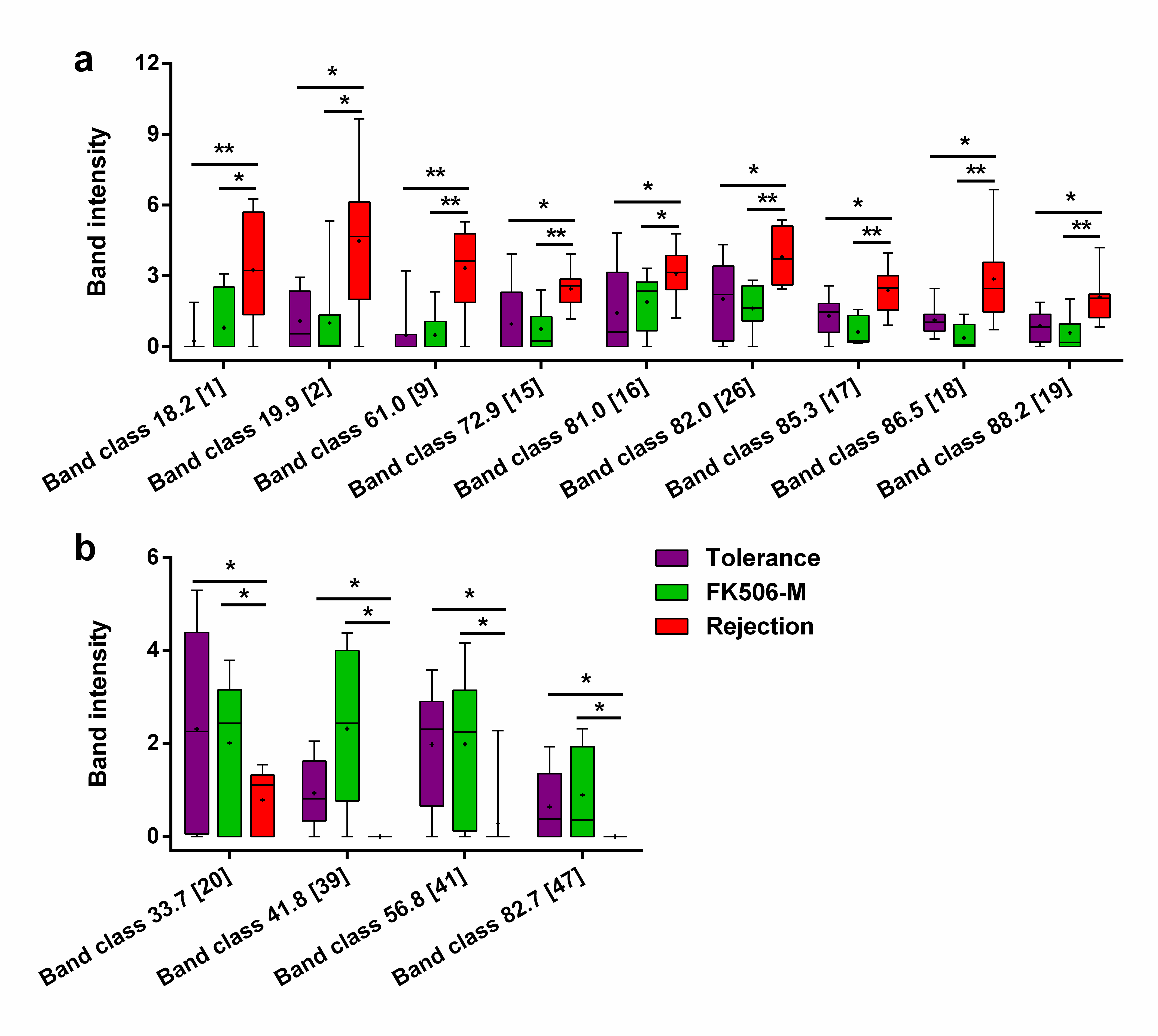
**Figure 2 An optimal dosage of FK506 maintained the gut barrier intact and resulted in low endotoxin levels after liver transplantation.**A: Representative intestinal mucosal ultrastructure shown by transmission electron microscopy for the different treatment groups (12500 ×, 30000 ×); B: Levels of plasma endotoxins in the different groups. M: Microvilli; TJ: Tight junction. a*P* < 0.05; b*P* < 0.01; c*P* < 0.001.

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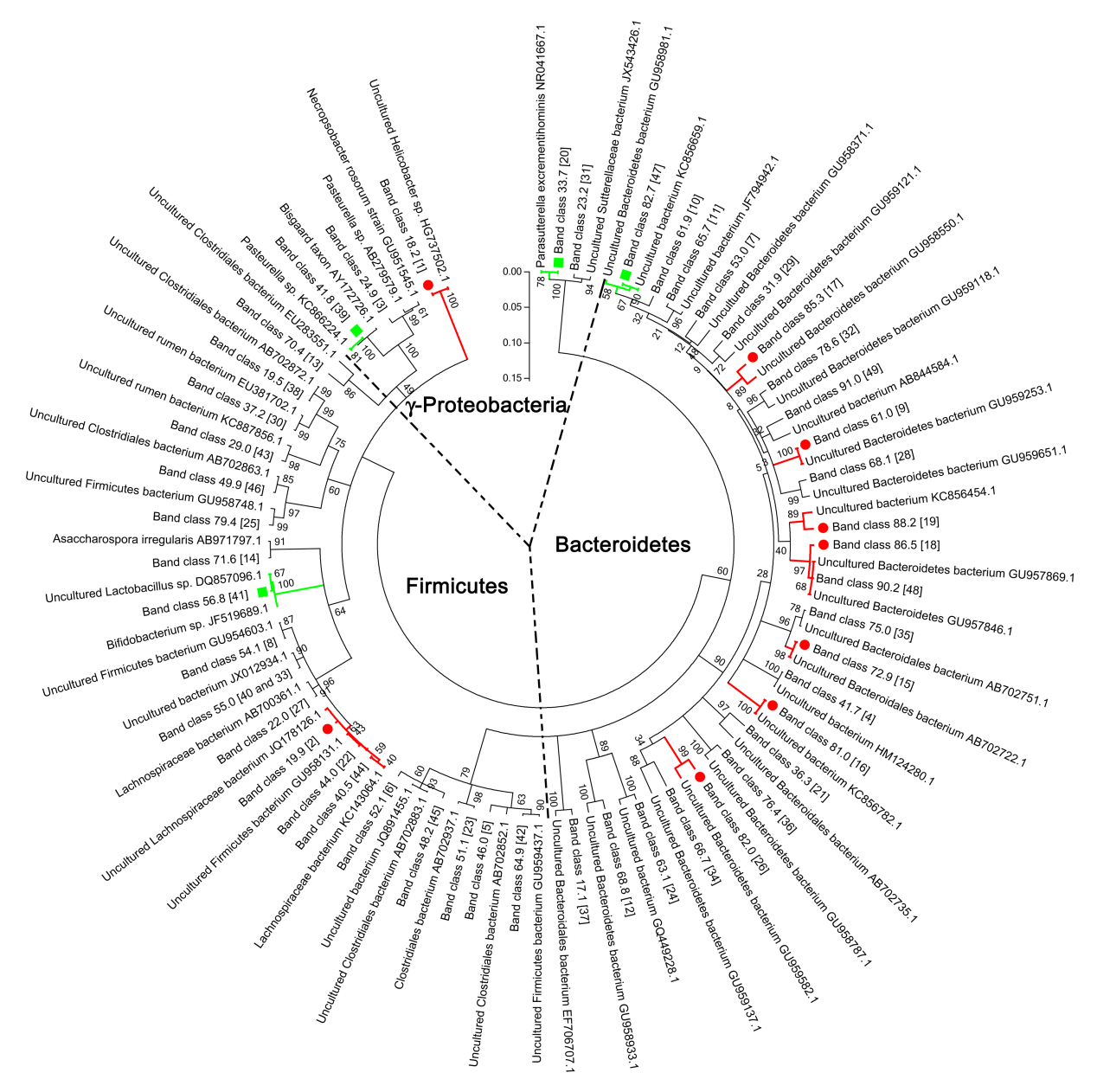
**Figure 3 Optimal dosage of FK506 induced a stable gut microbiota as determined using denaturing gradient gel electrophoresis.** Based on the DGGE method, the composition and distribution of ileocecal bacterial communities from samples obtained in rats following LT. The Dice similarity coefficient (band based) and unweighted pair-group method with arithmetic means (UPGMA) as a cluster method were utilized to analyze the similarity of each band in the different groups. Metric scale denotes the degree of similarity. Each band represents a bacterial clone. Band numbers indicated the position of bands excised for sequence analyses. DGGE: Denaturing gradient gel electrophoresis.

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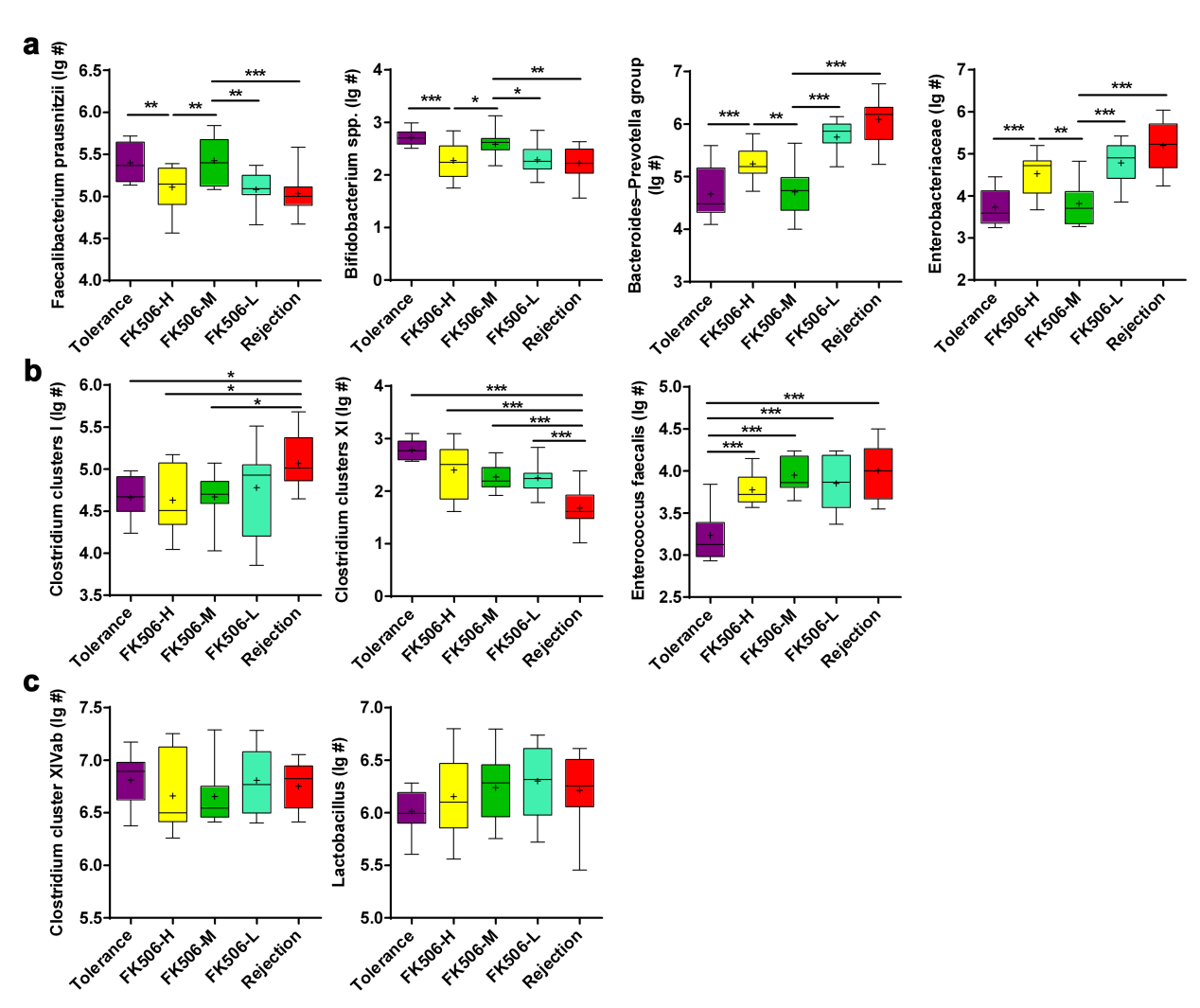
**Figure 4 Diversity analysis and cluster analysis of denaturing gradient gel electrophoresis profiles based on Dice’s coefficient and the unweighted pair-group method with arithmetic means (UPGMA) method.** A:Based on amount of gray area of each band of each lane in the denaturing gradient gel electrophoresis (DGGE) profiles quantified using the Gel-Pro analyzer, the gut microbial diversity and species richness were assessed with the Shannon index and Simpson index, in the different groups; B: The abundance and distribution of the rare species were estimated using the Chao-1 index, Fisher alpha index, and Menhinick index in the different groups; C: Multidimensional scaling (MDS) analysis of each sample of the DGGE profiles. The plot is an optimized three-dimensional representation of the similarity matrix obtained from BioNumerics software. The Euclidean distance between the two points reflects similarity; D: Principal components analysis (PCA) of fecal microbiota based on DGGE fingerprinting. The plot is orientated to maximize the variation among lanes along the first three principal components (with contributions of 13.7, 11.2 and 10.0, respectively) obtained from BioNumerics software. a*P* < 0.05; b*P* < 0.01; c*P* < 0.001.

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**Figure 5 Identification of crucial bands of gut microbial changes induced by FK506-M treatment. A:** Compared to the rejection group, the intensities of 9 band classes including 18.2 (band 1), 19.9 (band 2), 61.0 (band 9), 72.9 (band 15), 81.0 (band 16), 82.0 (band 26), 85.3 (band 17), 86.5 (band 18), and 88.2 (band 19), were significantly decreased in the FK506-M-treated and tolerance groups; B: Compared with the rejection group, 4 band classes including 33.7 (band 20), 41.8 (band 39), 56.8 (band 41), and 82.7 (band 47) showed increased intensities in the FK506-M-treated and tolerance groups. a*P* < 0.05, b*P* < 0.01.

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**Figure 6 Phylogenetic tree analysis of sequences in the denaturing gradient gel electrophoresis profiles**. Phylogenetic tree analysis of sequences from denaturing gradient gel electrophoresis (DGGE) profilesusing the neighbor-joining method was conducted using MEGA 5 software. Almost all matched bacteria of the 48 DGGE band classes were assigned to 3 phyla: *Bacteroidetes* (50.0%), *Firmicutes* (39.6%), and *Gammaproteobacteria* (10.4%). The fragment sequences were defined according to their positions in gels using the band-matching tool with BioNumerics software version 6.01 (Applied Maths). The numbers in the brackets were consistent with the numbers shown in DGGE profiling. Nine band classes (labeled with a red dot) were decreased and 4 band classes (labeled with the green square)were higher in the FK506-M group than the rejection group. The plot was obtained using MEGA5 software (<http://en.wikipedia.org/wiki/MEGA,_Molecular_Evolutionary_Genetics_Analysis>).

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**Figure 7 Quantitative verification of predominant bacterial community in fecal microbiota using RT-qPCR.** Gut dominant bacterial populations were analyzed with RT-qPCR for the different groups. The dominant bacteria mainly included *Faecalibacterium prausnitzii*, *Bifidobacterium spp.*, *Bacteroides-Prevotella* group, *Enterobacteriaceae*, *Clostridium clusters I*, *Clostridium clusters XI*, *Enterococcus faecalis*, *Clostridium cluster XIVab*, *Lactobacillus*. Statistical analysis was performed with one-way ANOVA. a*P* < 0.05, b*P* < 0.01, c*P* < 0.001. 1Log10 copies/g: log10 No. of 16S rDNA gene copies per gram feces (wet weight).