(54) HUMAN-DERIVED BACTERIA THAT INDUCE PROLIFERATION OR ACCUMULATION OF

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(58) Field of Classification Search
None
See application file for complete search history.

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(57) ABSTRACT

Species of human-derived bacteria belonging to the Clostridia class have been shown to induce accumulation of regulatory T cells (Treg cells) in the colon and suppress immune functions. Pharmaceutical compositions containing these bacteria can be used to prevent and treat immune-mediated diseases such as autoimmune diseases.

7 Claims, 22 Drawing Sheets
Specification includes a Sequence Listing.
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Fig. 1E

GF + huChloro
(#B4)

GF + huUT
(#A1)

GF

IL-17

CLPL

SLPL
Fig. 12

Fig. 13
Fig. 17

% Foxp3+ in CD4

GF + mix + 5-mix-A + 5-mix-B + 5-mix-C + 17-mix

ns

Fig. 18

% Foxp3+ in CD4

SPF + cont. SPF + 17-mix

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HUMAN-DERIVED BACTERIA THAT INDUCE PROLIFERATION OR ACCUMULATION OF
RELATED APPLICATIONS


TECHNICAL FIELD

The subject matter described herein relates to a composition of human-derived bacteria that induces proliferation, accumulation, or proliferation and accumulation of regulatory T cells and which comprises, as an active component, (a) one or more (a, at least one) human-derived bacteria that belongs to the Clostridia class, (b) culture supernatant of one or more (a, at least one) of the bacteria; (c) a physiologically active substance derived from one or more of the bacteria or (d) a combination of any two or more of the foregoing. It also relates to a method for inducing proliferation, accumulation or proliferation and accumulation of regulatory T cells. The composition, which comprises any of (a)-(d) above, is referred to as a bacterial composition. Moreover, the subject matter relates to a method for treating or preventing at least one disease or condition that is responsive to induction of regulatory T cells, such as autoimmune diseases, inflammatory diseases, and infectious diseases, by administering the bacterial composition to an individual in need thereof.

BACKGROUND

Hundreds of species of commensal microorganisms are harbored in the gastrointestinal tracts of mammals, where they interact with the host immune system. Research using germ-free (GF) animals has shown that the commensal microorganisms influence the development of the mucosal immune system, such as histogenesis of Peyer’s patches (PPs) and isolated lymphoid follicles (ILFs), secretion of antimicrobial peptides from the epithelium, and accumulation of unique lymphocytes in mucosal tissues, including immunoglobulin A-producing plasma cells, intraepithelial lymphocytes, IL-17-producing CD4-positive T cells (Th17), and IL-22-producing NK-like cells (Non-Patent Literature (NPL) 1 to 7). Consequently, the presence of intestinal bacteria enhances protective functions of the mucous membranes, enabling the host to mount robust immune responses against pathogenic microbes invading the body. On the other hand, the mucosal immune system maintains unresponsiveness to dietary antigens and harmless microbes (NPL Document 3). Abnormality in the regulation of cross-talk between commensal bacteria and the immune system (intestinal dysbiosis) may lead to overly robust immune response to environmental antigens and inflammatory bowel disease (IBD) may result (NPL 8 to 10).

Recent studies have shown that individual commensal bacteria control differentiation of their specific immune cells in the mucosal immune system. For example, Bacteroides fragilis, which is a commensal bacterium in humans, specifically induces a systemic Th1 cell response and a mucosal IL-10-producing T cell response in mice, and plays a role in protecting the host from colitis, which is caused by a pathogen (NPL 3). Segmented filamentous bacteria, which are intestinal commensal bacteria in mice, induce mucosal Th17 cell response and enhance resistance against infection of gastrointestinal tracts of the host with a pathogen (NPL 11 to 13). In addition, short-chain fatty acids derived from several commensal bacteria are known to suppress intestinal inflammation (NPL 14). Moreover, it has been observed that the presence of some species of intestinal microbiota greatly influences the differentiation of regulatory T cells (hereafter referred to as “Treg cells”) which help maintain homeostasis of the immune system. Although specific species of murine bacterial commensals that can strongly stimulate Tregs have been identified (NPL 15), it is still unknown whether species of human commensal bacteria exert an equivalent influence on the human immune system. Furthermore, the human intestinal tract harbors more than a thousand bacterial species, many of which have not yet been cultured (NPL 16). It is not feasible to guess a priori which ones, if any, might have an effect on Tregs.

In order to develop drugs, dietary supplements, or foods with beneficial immune functions for human use, it is desirable to identify commensal microorganisms that naturally colonize humans and have immune-modulating properties. Furthermore, since many of the commensals in the human microbiome have yet to be cultured, it is necessary to develop methods to cultivate them so that they can be produced by traditional industrial fermentation processes and subsequently incorporated in pharmaceutical or food formulations.

CD4⁺ T cells are regulatory T cells that have been identified as a cell subset that suppresses immunity. A transcription factor, Foxp3, is expressed in CD4⁺ T cells, which are known to play an important role in maintaining immunological homeostasis (NPL 8, 9, 17, and 18). Foxp3-expressing cells are present in large numbers in the colon and only Treg cells present locally in the colon constantly express IL-10, an immunosuppressive cytokine, at a high level (NPL 19). Animals having CD4⁺ Foxp3⁺ cells from which IL-10 is specifically removed develop inflammatory bowel disease (NPL 20).

Accordingly, there is a need to identify human-derived commensal bacterial species with the ability to strongly induce Treg cells to produce IL-10 in the colon at a high level and to develop methods to culture such species. Such species could be used to enhance immunosuppression, which, in turn, can be applied to treatment of autoimmune diseases, such as inflammatory bowel disease, inflammatory diseases, allergies, or organ transplantation, among other diseases and conditions.

NON PATENT LITERATURE

The present compositions and methods have been made in view of the above-described problems in the art. Described herein are methods of identifying and cultivating intestinal commensal bacteria, isolated from humans, which induce, preferably strongly, the proliferation, accumulation, or proliferation and accumulation of regulatory T cells. Described are compositions, also referred to as bacterial compositions, that (1) comprise (a) one or more of the identified intestinal commensal (human-derived) bacteria; (b) a culture supernatant of one or more of the bacteria; (c) one or more physiologically active substance derived from one or more of the bacteria or from one or more of the culture supernatant; (d) a combination of any two or three of (a)-(c) and (2) induce the proliferation and/or accumulation of regulatory T cells (Treg cells). Alternatively, a composition comprises (a) one or more of the identified intestinal commensal (human-derived) bacteria; (b) a culture supernatant of one or more of the bacteria; or (c) one or more physiologically active substance derived from the bacteria or from the culture supernatant, wherein the composition induces proliferation and/or accumulation of regulatory T cells. In some embodiments, the composition comprises one or more of the identified intestinal commensal (human-derived) bacteria. In some embodiments, the composition comprises a culture supernatant of one or more of the bacteria. In some embodiments, the composition comprises one or more physiologically active substance derived from the bacteria or from the culture supernatant. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is three or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is five or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is twenty-three or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is twenty-three or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is twenty-three or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is twenty-three or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is twenty-three or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is twenty-three or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is twenty-three or more.
nospiraceae bacterium 5_1_57FAA, Lachnospiraceae bacterium 6_1_63FAA, Clostridium botulinum ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes caccae, Clostridium botulinum, Lachnospiraceae bacterium 3_1_57FAA, CT1, Anaerotruncus colihominis, Anaerotruncus colihominis DSM 17241, Ruminococcus sp. ID8, Lachnospiraceae bacterium 2_1_46FAA, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum WAL-14163, Eubacterium contortum, Clostridium sp. D5, Oscillosporae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47FAA, Blautia cocoides, Anaerostipes caccae DSM 14662. In some embodiments, a bacterial composition comprises a culture supernatant of at least one (a, one or more) of the bacteria described/listed herein. In some embodiments, a bacterial composition comprises a physiologically active substance derived from (a, one or more) of the bacteria described/listed herein. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is three or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is five or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is 17 or more. In some embodiments, the one or more bacteria of one or more physiologically active substance derived from the bacteria is 23 or more. In some embodiments, the one or more bacteria or one or more physiologically active substance derived from the bacteria is 23. Bacterial compositions can comprise any bacteria (Clostridia or other bacteria) that contain DNA comprising a nucleotide sequence having sufficient homology with sequences provided herein and that exhibit substantially the same effect on regulatory T cells as that exerted by any one of the following: Clostridium saccharogena, Clostridium ramsonum JCM1298, Clostridium ramsonum, Flavonifractor plautii, Pseudoflavonifractor capsulatus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57FAA, Lachnospiraceae bacterium 6_1_63FAA, Clostridium sp. 14616, Clostridium botulinum ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes caccae, Clostridium botulinum, Lachnospiraceae bacterium 3_1_57FAA, CT1, Anaerotruncus colihominis, Anaerotruncus colihominis DSM 17241, Ruminococcus sp. ID8, Lachnospiraceae bacterium 2_1_46FAA, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum WAL-14163, Eubacterium contortum, Clostridium sp. D5, Oscillosporae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47FAA, Blautia cocoides, Anaerostipes caccae DSM 14662. In specific embodiments, bacterial compositions comprise a bacterial composition that contain DNA comprising a nucleotide sequence having at least 97%, 98% or 99% homology with sequences provided herein, such as, but not limited to, the nucleotide sequences designated OTU herein and listed, for example, at the pages following the last Example. In specific embodiments, the bacteria contain DNA comprising a nucleotide sequence that has at least 90% (90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or 100%) homology with one or more DNA sequence designated herein as follows: OTU136; OTU46; OTU221; OTU9; OTU296; OTU21; OTU166; OTU73; OTU174; OTU14; OTU55; OTU337; OTU193; OTU15; OTU206; OTU87; OTU86; OTU152; OTU253; OTU258; OTU281; OTU288; OTU334; OTU359; OTU362; or OTU367. Alternatively, the bacteria contain DNA comprising a nucleotide sequence that has at least 90% (90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or 100%) homology with DNA of one or more of the following: Clostridium saccharogena, Clostridium ramsonum JCM1298, Clostridium ramsonum, Flavonifractor plautii, Pseudoflavonifractor capsulatus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57FAA, Lachnospiraceae bacterium 6_1_63FAA, Clostridium botulinum ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes caccae, Clostridium botulinum, Lachnospiraceae bacterium 3_1_57FAA, CT1, Anaerotruncus colihominis, Anaerotruncus colihominis DSM 17241, Ruminococcus sp. ID8, Lachnospiraceae bacterium 2_1_46FAA, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum WAL-14163, Eubacterium contortum, Clostridium sp. D5, Oscillosporae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47FAA, Blautia cocoides, Anaerostipes caccae DSM 14662.
Oscillatoria valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1.7.47FAA, Blautia cocoideae, Anaerostipes caccae DSM 14662. Any of the bacteria of the Clostridia class can be present in spore form or vegetative form.

Solution of Problem

As described herein, among the more than a thousand species of bacteria in the human microbiome, there are several species that strongly induce the accumulation of Tregs in the colon. As also described, although most bacterial species present in fecal samples from healthy individuals do not have the ability to stimulate Tregs, species that belong to the Clostridia class have the ability to cause a robust induction of Tregs in the colon. Moreover, the inventors have obtained in vitro cultures of each of the bacterial species identified and shown that inoculating mice with the in vitro cultured species also leads to a robust accumulation of Tregs in the colon.

As described herein, compositions that comprise, as an active component, (a) one or more of certain species of bacteria that belong to the Clostridia class or bacteria that contain DNA comprising a nucleotide sequence having at least 90% homology with sequences provided herein, in spore form or in vegetative form; (b) a culture supernatant of one or more such bacteria; (c) one or more physiologically active substance derived from (a) or (b); or (d) a combination of any two or three of (a), (b), and (c) and induce the proliferation and/or accumulation of regulatory T cells (Treg cells) suppress immune functions.

More specifically:

One embodiment is a composition that induces proliferation, accumulation or both proliferation and accumulation of regulatory T cells, the composition comprising, as an active component, at least one organism and/or at least one substance selected from the group consisting of: Clostridium saccharogenum, Clostridium ramsonum JCM1298, Clostridium ramsonum, Flavonifractor plautii, Pseudoflavonifactor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5.1.57FAA, Lachnospiraceae bacterium 6.1.63FAA, Clostridium sp. 14616, Clostridium botelae ATCC BAA-613, cf. Clostridium sp. MALG05, Erysipelotrichaceae bacterium 2.2.44A, Clostridium indolios, Anaerostipes cacaee, Clostridium botelae, Clostridiales bacterium DIF_VP50, Lachnospiraceae bacterium 3.1.57FAA_CT1, Anaerotranscus coliforminis, Anaerotranscus coliforminis DSM 17241, Ruminococcus sp. D8B, Lachnospiraceae bacterium 2.1.46FAA, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosus, Clostridium symbiosus WAL-14163, Eubacterium contortum, Clostridium sp. D5, Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1.7.47FAA, Blautia cocoideae, Anaerostipes cacaee DSM 14662; a culture supernatant of at least one of the bacteria described/listed herein, and a physiologically active substance derived from a bacterium described/listed herein.

In some embodiments, the active component is one or more of Clostridium saccharogenum, Clostridium ramsonum JCM1298, Clostridium ramsonum, Flavonifractor plautii, Pseudoflavonifactor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5.1.57FAA, Lachnospiraceae bacterium 6.1.63FAA, Clostridium sp. 14616, Clostridium botelae ATCC BAA-613, cf. Clostridium sp. MALG05, Erysipelotrichaceae bacterium 2.2.44A, Clostridium indolios, Anaerostipes cacaee, Clostridium botelae, Clostridiales bacterium DIF_VP50, Lachnospiraceae bacterium 3.1.57FAA_CT1, Anaerotranscus coliforminis, Anaerotranscus coliforminis DSM 17241, Ruminococcus sp. D8B, Lachnospiraceae bacterium 2.1.46FAA, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosus, Clostridium symbiosus WAL-14163, Eubacterium contortum, Clostridium sp. D5, Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1.7.47FAA, Blautia cocoideae, Anaerostipes cacaee DSM 14662 or any bacteria (such as human-derived bacteria) that contain DNA comprising at least 90% homol-
ogy (e.g., 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or 100% homology) with sequences provided herein, such as, but not limited to, the nucleotide sequences designated OTU herein and listed, for example, at the pages following the last Example. In specific embodiments, the bacteria contain DNA comprising a nucleotide sequence that has at least 97%, at least 98% or at least 99% homology with one or more DNA sequence designated herein as follows: OTU136; OTU46; OTU221; OTU9; OTU296; OTU21; OTU166; OTU73; OTU174; OTU14; OTU55; OTU337; OTU314; OTU195; OTU306; OTU87; OTU86; OTU1152; OTU253; OTU259; OTU281; OTU288; OTU334; OTU359; OTU362; or OTU367. Alternatively, the bacteria contain DNA comprising a nucleotide sequence that has at least 97% (97%, 98%, 99%, 100%) homology with DNA of one or more of the following: Clostridium saccharogenum, Clostridium ramosum JCM1298, Clostridium ramosum, Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57FAA, Lachnospiraceae bacterium 6_1_63FAA, Clostridium bolteae ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes cacciae, Clostridium bolteae, Lachnospiraceae bacterium 3_1_57FAA, CT1, Anaerotruncus colihominis, Anaerotruncus colihominis DSM 17241, Ruminococcus sp. ID8, Lachnospiraceae bacterium 2_1_46FAA, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum WAL-14163, Eubacterium centorum, Clostridium sp. D5, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47FAA, Blautia coelioides, and Anaerostipes cacciae DSM 14662.

In one embodiment, the composition induces regulatory T cells that are transcription factor Foxp3-positive regulatory T cells or IL-10-producing regulatory T cells. In another embodiment, the composition has an immunosuppressive effect.

One embodiment is a pharmaceutical composition that induces proliferation, accumulation or both proliferation and/or accumulation of regulatory T cells and suppresses immune function. The pharmaceutical composition comprises a bacterial composition described herein and a pharmaceutically acceptable component, such as a carrier, a solvent or a diluent. In specific embodiments, such a pharmaceutical composition comprises (a) one or more species of bacteria belonging to the Clostridia class, as described herein, in spore form or in vegetative form, (b) a culture supernatant of such bacteria, (c) a physiologically active substance derived therefrom or (d) a combination of any two or three of (1), (2) and (3) and (b) a pharmaceutically acceptable component, such as a carrier, a solvent or a diluent. In specific embodiments, (a) above is at least one organism or substance selected from the group consisting of: Clostridium saccharogenum, Clostridium ramosum JCM1298, Clostridium ramosum, Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57FAA, Lachnospiraceae bacterium 6_1_63FAA, Clostridium bolteae ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes cacciae, Clostridium bolteae, Lachnospiraceae bacterium 3_1_57FAA, CT1, Anaerotruncus colihominis, Anaerotruncus colihominis DSM 17241, Ruminococcus sp. ID8, Lachnospiraceae bacterium 2_1_46FAA, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum WAL-14163, Eubacterium centorum, Clostridium sp. D5, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47FAA, Blautia coelioides, and Anaerostipes cacciae DSM 14662. In some embodiments, (1) above is a culture supernatant of one or more of the bacteria. In some embodiments, (a) above is a physiologically active substance derived from one or more of the bacteria. In some embodiments, the at least one organism or substances is three or more. In some embodiments, the at least one organism or substances is five or more. In some embodiments, the at least one organism or substances is 17 or more. In some embodiments, the at least one organism or substances is 23 or more. In some embodiments, the at least one organism or substances is 23. In further embodiments, (a) above is bacteria (such as human-derived bacteria) that contain DNA comprising at least 90% homology (e.g., 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or 100% homology) with sequences provided herein, such as, but not limited to, the nucleotide sequences designated OTU herein and listed, for example, at the pages following the last Example. In specific embodiments of the pharmaceutical composition, the bacteria contain DNA comprising a nucleotide sequence that has at least 97%, at least 98% or at least 99% homology with one or more DNA sequence designated herein as follows: OTU136; OTU46; OTU221; OTU9; OTU296; OTU21; OTU166; OTU73; OTU174; OTU14; OTU55; OTU337; OTU314; OTU195; OTU306; OTU87; OTU86; OTU1152; OTU253; OTU259; OTU281; OTU288; OTU334; OTU359; OTU362; or OTU367. Alternatively, the bacteria in the pharmaceutical composition contain DNA comprising a nucleotide sequence that has at least 97% (97%, 98%, 99%, 100%) homology with DNA of one or more of the following: Clostridium saccharogenum,
Clostridium ramosum JCM1298, Clostridium ramosum, Flavonifractor plautii, Pseudo flavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium sacccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57FAA, Lachnospiraceae bacterium 6_1_63FAA, Clostridium sp. 14616, Clostridium bolteae ATCC BAA-613, clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes caccae, Clostridium bolteae, Lachnospiraceae bacterium D3F_VP30, Lachnospiraceae bacterium 3_1_57FAA_CTI, Anaerotruncus coliformis, Anaerotruncus coliformis D3F_VP30, Ruminococcus sp. IDB, Lachnospiraceae bacterium 2_1_46FAA, Clostridium lavalense, Clostridium asparagiforme D3F_VP30, Clostridium symbiosum, Clostridium symbiosum WAL-14163, Eubacterium contortum, Clostridium sp. D5, Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47FAA, Blautia cocoides, Anaerostipes caccae D3F_VP30.

The pharmaceutical composition induces the proliferation and/or accumulation of regulatory T cells (Tregs) and suppresses immune function.

Also provided is a method of inducing proliferation, accumulation or both proliferation and accumulation of regulatory T cells in an individual (e.g., an individual in need thereof, such as an individual in need of induction of proliferation and/or accumulation of regulatory T cells). The method comprises administering to the individual a bacterial composition described herein or a pharmaceutical composition comprising a bacterial composition described herein. In the method at least one organism or substance selected from the group consisting of: Clostridium saccharogroupon, Clostridium ramosum JCM1298, Clostridium ramosum, Flavonifractor plautii, Pseudo flavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57FAA, Lachnospiraceae bacterium 6_1_63FAA, Clostridium sp. 14616, Clostridium bolteae ATCC BAA-613, clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes caccae, Clostridium bolteae, Lachnospiraceae bacterium D3F_VP30, Lachnospiraceae bacterium 3_1_57FAA_CTI, Anaerotruncus coliformis, Anaerotruncus coliformis D3F_VP30, Ruminococcus sp. IDB, Lachnospiraceae bacterium 2_1_46FAA, Clostridium lavalense, Clostridium asparagiforme D3F_VP30, Clostridium symbiosum, Clostridium symbiosum WAL-14163, Eubacterium contortum, Clostridium sp. D5, Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47FAA, Blautia cocoides, Anaerostipes caccae D3F_VP30.

Also provided herein is a composition that comprises the bacterial composition and at least one prebiotic substance.

The bacterial composition may be administered in combination with a substance selected from the group consisting of corticosteroids, mesalazine, mesalamine, sulfasalazine, sulfasalazine derivatives, immuno suppressive drugs, cyclosporin A, mercaptopurine, azathiopurine, prednisone, methotrexate, anti-arthimatic, glucocorticoids, epinephrine, theophylline, cromolyn sodium, anti-leukotrienes, anti-cholinergic drugs for rhinitis, anti-cholinergic decongestants, mast-cell stabilizers, monoclonal anti-IgE antibodies, vaccines, anti-TNF inhibitors such as infliximab, adalimumab, certolizumab pegol, golimumab, or etanercept, and combinations thereof. Also described herein is a composition that comprises the bacterial composition and at least one substance selected from the group consisting of corticosteroids, mesalazine, mesalamine, sulfasalazine, sulfasalazine derivatives, immuno suppressive drugs, cyclosporin A, mercaptopurine, azathiopurine, prednisone, methotrexate, anti-arthimatic, glucocorticoids, epinephrine, theophylline, cromolyn sodium, anti-leukotrienes, anti-cholinergic drugs for rhinitis, anti-cholinergic decongestants, mast-cell stabilizers, monoclonal anti-IgE antibodies, vaccines, anti-TNF inhibitors such as infliximab, adalimumab, certolizumab pegol, golimumab, or etanercept, and combinations thereof.

In a further embodiment, the bacterial composition can be used as an adjuvant to improve the efficacy of a vaccine formulation. For example, the bacterial composition can be used as an adjuvant to a vaccine for the prophylaxis or treatment of an autoimmune disease or an allergic disease. In some embodiments, a method for prophylaxis or treatment is provided, the method comprising administering the bacterial composition and administering a vaccine.

Assessment of the extent of induction of proliferation or accumulation of regulatory T cells that results from administration of a composition described herein can be carried out by a variety of approaches, such as by measurement of the number of Foxp3-expressing Tregs in a patient sample (such as a biopsy or a blood sample), promotion of IL-10 expression, promotion of CTLA4 expression, promotion of IDO expression, suppression of IL-4 expression, or colonization of an individual with the bacterial composition. The results of such assessments are used as an index of the induction of proliferation or accumulation of regulatory T cells in the individual.

In one embodiment, administration of a composition described herein causes induction of the regulatory T cells that are transcription factor Foxp3-positive regulatory T cells or IL-10-producing regulatory T cells.

The composition described herein can be administered by a variety of routes and in one embodiment, is administered orally to an individual in need thereof, such as a patient in
need thereof. The composition may be administered in a number of oral forms, such as in a powder or dissolved in a liquid formulation), in enteric capsules, in sachets, or in a food matrix, such as yogurt, or a drink.

Also provided is a method to predict a subject’s response to treatment (predict whether the subject will respond to treatment) with compositions of the invention. The method comprises (a) obtaining a (at least one, one or more) sample, such as a fecal sample or a colonic biopsy, from a patient before he or she is treated with a bacterial composition described herein; (b) measuring or determining the percentage or absolute counts in the sample of at least one bacterial species selected from the group consisting of: Clostridium saccharogumnia, Clostridium ramosum JCM1298, Clostridium ramosum, Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57F, Lachnospiraceae bacterium 6_1_63F, Clostridium sp. 14616, Clostridium botulinum ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes cacae, Clostridium botulinum, Lachnospiraceae bacterium DJF_VP30, Lachnospiraceae bacterium 3_1_57F AA_CT1, Anaerotruncus colinomis, Anaerotruncus colinomis DSM 17241, Ruminococcus sp. IDB, Lachnospiraceae bacterium 2_1_46F, Clostridium lavalense, Clostridium asparagusiforme DSM 15981, Clostridium symbiosum, Clostridium ramosum JCM1298, Eubacterium contortum, Clostridium sp. D5, Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002, and Clostridiales bacterium 1_7_47F, Blautia cacae, and Anaerostipes cacae DSM 14662, thereby producing a percentage or count, and (c) comparing the resulting percentage or count (measurement) to a baseline value of the same measurement in a healthy subject, wherein a percentage or count in the sample obtained from the patient that is lower than the baseline value indicates that the subject may respond favorably to administration of the bacterial composition. In some embodiments, the method further comprises (d) administering the bacterial composition to the patient if the percentage or count in the sample obtained from the patient is lower than the baseline value. Optionally, the method may further comprise measuring in a patient’s sample (e.g., a fecal sample or a colonic biopsy) the percentages or absolute counts of other commensal species that belong to Clostridium Clusters IV and XIVA, but are not present in the bacterial composition, wherein a value of the percentage or absolute count (measurement) lower than baseline further indicates that the subject may respond favorably to administration of the bacterial compositions. In some embodiments, the method further comprises administering the bacterial composition to the patient if the value of the percentage or absolute count (measurement) is lower than baseline. In one embodiment, the patient being assessed suffers from inflammatory bowel disease or a Clostridium difficile infection.

Also provided is a method of monitoring a subject's response to treatment with the bacterial compositions of the invention, comprising: (a) obtaining a (at least one) sample, such as a fecal sample or a colonic biopsy from a patient before treatment with a bacterial composition described herein; (b) obtaining, a (at least one) corresponding sample from the patient after treatment with a bacterial composition described herein; and (c) comparing the percentage or absolute counts of at least one bacterial species selected from the group consisting of: Clostridium saccharogumnia, Clostridium ramosum JCM1298, Clostridium ramosum, Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57F, Lachnospiraceae bacterium 6_1_63F, Clostridium sp. 14616, Clostridium botulinum ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes cacae, Clostridium botulinum, Lachnospiraceae bacterium DJF_VP30, Lachnospiraceae bacterium 3_1_57F AA_CT1, Anaerotruncus colinomis, Anaerotruncus colinomis DSM 17241, Ruminococcus sp. IDB, Lachnospiraceae bacterium 2_1_46F, Clostridium lavalense, Clostridium asparagusiforme DSM 15981, Clostridium symbiosum, Clostridium ramosum JCM1298, Eubacterium contortum, Clostridium sp. D5, Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002, and Clostridiales bacterium 1_7_47F, Blautia cacae, Anaerostipes cacae DSM 14662 in the sample obtained in (a) with the percentage or absolute counts of the same at least one bacterial species in the sample obtained in (b), wherein a higher value in the sample obtained in (b) (after treatment with the bacterial composition) than in the sample obtained in (a) (before treatment) indicates that the subject has responded favorably to treatment (e.g., is a positive indicator of enhanced immunosuppression in the subject). In some embodiments, the method further comprises (d) further administering the bacterial composition to the patient or ceasing administration of the bacterial composition to the patient based on the comparison in (c). Optionally, the method may further comprise measuring in the subject’s samples the percentages or absolute counts of other commensal species that belong to Clostridium Clusters IV and XIVA, but are not present in the bacterial composition, wherein a higher value after treatment than before treatment indicates that the subject has responded favorably to treatment.

Effects of Compositions and Methods Described Herein

The compositions described herein, which contain, as an active component, selected bacteria belonging to the Clostridia class or other bacteria, as described herein; a culture supernatant of such bacteria; a physiologically active substance derived from such bacteria; or a combination of two or three of the foregoing are excellent at inducing the proliferation or accumulation of regulatory T cells (Treg cells).

Immunity in an individual can be suppressed through administration of the subject composition, such as through ingestion of the bacterial composition in a food or beverage or as a dietary supplement or through administration of a pharmaceutical composition comprising the bacterial composition. The subject composition can be used, for example, to prevent or treat autoimmune diseases, allergic diseases, infectious diseases, as well as to suppress immunological rejection in organ transplantation or the like. In addition, if a food or beverage, such as a health food, comprises the subject composition, healthy individuals can ingest the composition easily and routinely. As a result, it is possible to induce the proliferation and/or accumulation of regulatory T cells and thereby to improve immune functions.

The composition described herein provides for a natural, long-lasting, patient-friendly, and benign treatment alternative for immune-mediated conditions. For example, inflam-
BRIEF DESCRIPTION OF DRAWINGS

FIG. 1A is a histogram showing Foxp3 expression gated CD4+ cells from colonic lamina propria (C LPL, left panel) and small intestinal lamina propria (SI LPL, right panel) of GF mice or GF mice colonized with untreated (+huUT, n=4, numbering from #A1 to #A4) or chloroform-treated (+huChloro, n=4, numbering from #B1 to #B4) human feces. Numbers above bracketed lines indicate the percentage of the population.

FIG. 1B is a histogram showing Helios expression in Foxp3+CD4+ cells from colonic lamina propria (left panel) and small intestinal lamina propria (right panel) of GF mice or GF mice colonized with untreated (+huUT) or chloroform-treated (+huChloro) human feces. Numbers above bracketed lines indicate the percentage of the population.

FIGS. 1C-1D are graphs showing, respectively, combined data for Foxp3 expression in CD4+ cells, and for Helios expression in Foxp3+CD4+ cells, from colonic lamina propria (left panel) and small intestinal lamina propria (right panel) of GF mice or GF mice colonized with untreated (+huUT) or chloroform-treated (+huChloro) human feces. Each circle in FIG. 1C represents a separate animal, and error bars indicate the SD. *P<0.05; **P<0.001, unpaired t test.

FIG. 1E shows representative flow cytometry dot plots for the intracellular expressions of IL-17 and IFN-γ in CD4+ cells from colonic lamina propria (upper panel) and small intestinal lamina propria (lower panel) of GF mice or GF mice colonized with untreated (+huUT) or chloroform-treated (+huChloro) human feces. The number in each quadrant indicates the percentage of the population.

FIGS. 1F-1G show, respectively, combined data of all mice for IL-17 and IFN-γ expression in CD4+ cells from colonic lamina propria (left panel) and small intestinal lamina propria (right panel) of GF mice or GF mice colonized with untreated (+huUT) or chloroform-treated (+huChloro) human feces. Each circle in FIG. 1F and FIG. 1G represents a separate animal, and error bars indicate the SD. *P<0.05; ns, not significant (P>0.05), unpaired t test.

FIGS. 2A-2C show representative plots (FIG. 2A) and combined data (FIGS. 2B-2C) for Foxp3 expression in CD4+ cells (upper panel in FIG. 2A, left panel in FIG. 2B), or Helios expression in Foxp3+CD4+ cells (lower panel in FIG. 2A, right panel in FIG. 2C) for GF mice and GF mice orally inoculated (once a week for 4 weeks) with a suspension of chloroform-treated human feces that had been previously autoclaved. Numbers above bracketed lines in FIG. 2A indicate the percentage of the population. Each circle in FIG. 2B and FIG. 2C represents a separate animal, and error bars indicate the SD. *P<0.05, unpaired t test.

FIGS. 3A-3B show representative plots (FIG. 3A, data of mouse #C4 is shown here) and combined data (FIG. 3B) for Foxp3 expression in CD4+ cells from colonic and small intestinal lamina propria lymphocytes for GF mice and GF mice orally inoculated with chloroform-treated human feces (+huChloro, n=7, numbering from #C1 to #C7). Numbers above bracketed lines in FIG. 3A indicate the percentage of the population. Each circle in FIG. 3B represents a separate animal, and error bars indicate the SD. **P<0.001, unpaired t test.

FIGS. 4A-4B show representative plots (FIG. 4A) and combined data (FIG. 4B) for Foxp3 expression in CD4+ cells from colonic lamina propria (C LPL) and small intestinal lamina propria (SI LPL) for GF mice and GF mice (numbering from #D1 to #D6) that were co-housed with #C6 and #C7 ex-GF mice colonized with chloroform-treated human feces. Numbers above bracketed lines in FIG. 4A indicate the percentage of the population. Each circle in FIG. 4B represents a separate animal, and error bars indicate the SD. **P<0.001, unpaired t test.

FIGS. 5A-5C show representative plots and combined data for Foxp3 expression in CD4+ cells (FIG. 5A, 5B), or Helios expression in Foxp3+CD4+ cells (FIG. 5C) from colonic lamina propria (C LPL) and small intestinal lamina propria (SI LPL) for GF mice and GF mice that were inoculated with 2000-fold (+2000) n=4, numbering from #E1 to #E4) or 20000-fold (+20000, n=8, numbering from #F1 to #F8) diluted fecal suspension from #C4 mouse. Numbers above bracketed lines in FIG. 5A indicate the percentage of the population. Each circle in FIG. 5B and FIG. 5C represents a separate animal, and error bars indicate the SD. *P<0.05; **P<0.001, unpaired t test.

FIGS. 6A-6D show representative plots (FIGS. 6A, 6B) and combined data (FIGS. 6C, 6D) for Foxp3 expression in CD4+ cells (FIG. 6A, 6C), or Helios expression in Foxp3+CD4+ cells (FIGS. 6B, 6D) from colonic lamina propria (C LPL) and small intestinal lamina propria (SI LPL) for GF mice and GF mice that were inoculated with fecal suspension of #F3 (n=5), #F7 (n=4) or #F8 (n=4) mouse. Numbers above bracketed lines in FIG. 6A and FIG. 6B indicate the percentage of the population. Each circle in FIG. 6C and FIG. 6D represents a separate animal, and error bars indicate the SD. *P<0.05, **P<0.001, unpaired t test.

FIGS. 7A-7C show representative plots (FIG. 7A) and combined data (FIGS. 7B, 7C) for Foxp3 expression in CD4+ cells (FIGS. 7A, 7B) or Helios expression in Foxp3+CD4+ cells for GF mice and GF mice that were inoculated with 3 isolated strains of bacteria from cecal content of #F8 mouse (n=4, numbering from #J1 to #J4). Numbers above bracketed lines in FIG. 7A indicate the percentage of the population. Each circle in FIG. 7B and FIG. 7C represents a separate animal, and error bars indicate the SD. ns, not significant (P>0.05), unpaired t test.

FIG. 8 shows the relative abundances of OTUs having the same closest relative in each cecal sample (bacterial DNA was extracted from the cecal contents of mouse #A1, #C4, #F8, #G2, #H5, #J3, #J3 and #K3, shown in the bars). Total number of OTUs detected in each sample is depicted below the bar. The detected OTU names in sample #H3, #J3 or #K3, their closest relative and their similarity with the closest relative are depicted in the right table.

FIGS. 9A-9C show representative plots (FIG. 9A) and combined data (FIG. 9B, 9C) for Foxp3 expression in CD4+ cells (FIG. 9A, 9B), or Helios expression in Foxp3+CD4+ cells (FIG. 9A, 9C) from colonic lamina propria (C LPL) and small intestinal lamina propria (SI LPL) for GF mice and GF mice that were inoculated with bacteria collections from culture plate of cecal content of #G2 mouse (n=4, numbering from #K1 to #K4). Numbers above bracketed lines in FIG. 9A indicate the percentage of the population. Each circle in FIG. 9B and FIG. 9C represents a separate animal, and error bars indicate the SD. *P<0.05, **P<0.001, unpaired t test.

FIG. 10 shows the relative abundances of OTUs having the same closest relative in each cecal sample (bacterial DNA was extracted from the cecal contents of mouse #A1, #C4, #F8, #G2, #H5, #J3, #J3 and #K3, shown in the bars). Total number of OTUs detected in each sample is depicted below the bar. The detected OTU names in sample #H3, #J3 or #K3, their closest relative and their similarity with the closest relative are depicted in the right table.
FIGS. 10A-10C shows representative plots (FIG. 10A) and combined data (FIGS. 10B, 10C) for Foxp3 expression in CD4+ cells (FIG. 10A, 10B), or Helios expression in Foxp3+CD4+ cells (FIG. 10A) from colonic lamina propria (C LP) and small intestinal lamina propria (SI LP) for GF mice and GF mice that were inoculated with a mixture of 23 bacterial strains that were isolated and shown in Table 2 (23 mix). Numbers above bracketed lines in FIG. 10A indicate the percentage of the population. Each circle in FIG. 10B and FIG. 10C represent a separate animal, and error bars indicate the SD. *P<0.05; **P<0.001, unpaired t test.

FIG. 11 shows a representative plot of the accumulation of Foxp3+CD4+ cells in adult GF mice that were inoculated with 2×10^6 to 2×10^7-fold diluted caecal samples from +hungChio mice. Experiments were performed more than twice. Error bars indicate SD. **P<0.01, *P<0.05, as calculated by Student’s t-test.

FIG. 12 shows a representative plot of the accumulation of Foxp3+CD4+ cells in the colon of adult GF mice that were inoculated with a mixture of 23 bacterial strains that were isolated and shown in Table 2 (23-mix), chloroform-treated human feces (+hungChio) and Faecalibacterium prausnitzii (+Faecalii). Error bars indicate SD. **P<0.01, as calculated by Student’s t-test.

FIG. 13 shows a representative plot of the accumulation of Foxp3+CD4+ cells in adult GF mice that were the secondary (2×10^9-re) and tertiary (2×10^9-re) recipients of inoculations with the caecal content of 4×10^9 mice, and adult GF mice inoculated with 2×10^7-fold diluted caecal samples from 2×10^9 mice (+2×10^9).

FIG. 14 shows the results of 16s rDNA pyrosequencing the caecal contents from the defined mice (+hungChio, +hungChio, +2×10^4, +2×10^5, +2×10^5, +2×10^5, +2×10^5) using a 454 sequencer. The relative abundance of OTUs (%) in the caecal bacterial community in each mouse and the closest strains in the database and the corresponding isolated strain number for the indicated OTUs are shown.

FIG. 15 shows a representative plot of the accumulation of Foxp3+CD4+ cells in the colon of adult IqI BALB and B6 GF mice on inoculation with a mixture of 17 bacterial strains that were isolated and shown in Table 4 (17-mix), **P<0.01, as calculated by Student’s t-test.

FIG. 16 shows a representative plot of the accumulation of Foxp3+CD4+ cells in adult IqI GF mice mono-colonized with each of the 17 strains listed in Table 4 (17-mix).

FIG. 17 shows a representative plot of the accumulation of Foxp3+CD4+ cells in adult IqI GF mice colonized with 3-mix, 5 mix-A, 5 mix-B, 5 mix-C or 17 mix as listed in Table 4. Circles indicate individual animals. Experiments were performed more than twice with similar results. Error bars indicate SD. **P<0.01, *P<0.05, ns, not significant, as calculated by Student’s t-test.

FIG. 18 shows a representative plot of the accumulation of Foxp3+CD4+ cells in adult SPF mice repeatedly inoculated with 17 mix (SPF+17 mix; n=5) or control (SPF+control; n=6). **P<0.001, as calculated by Student’s t-test.

FIG. 19 shows the effects of inoculation with 17 mix on an OVA model of diarrhea, as measured by a quantitative diarrhea score. *P<0.05, as calculated by Student’s t-test.

FIG. 20 shows the survival of adult mice inoculated with a mixture of 17 bacterial strains listed in Table 4 (17-mix) following exposure to trimethoprim sulfonamide (TNBS), an agent used in experimental models of colitis.

FIG. 21 shows the relative abundance of each of the 17-mix strains in the human fecal microbiota of ulcerative colitis and healthy subjects. The publically available reads of 15 healthy and 20 ulcerative colitis subjects in the MetaHIT database were aligned to the genome of the 17 strains. The mean numbers of mapped reads in healthy and UC groups for each of the 17 strain genomes are shown. Error bars represent SEM. *P<0.05, as calculated by the Student’s t-test.

Table 1 shows the numbers of detected reads and the closest relatives for each OTU obtained from classification of sequences (3400 reads for each sample) resulting from 16s rDNA coding gene amplification and PCR metasequencing of bacterial DNA extracted from the caecal contents of mouse #A1, #B1, #F8, #G2, #H3, #I2, #J3 and #K3 (classification on the basis of sequence similarity, >97% identity to the sequences in nucleic acid databases using BLAST).

Table 2 shows, for each of seventeen bacterial strains isolated from the caecal contents of mouse #F8, #G2, #H1 and #K3 using BL agar or EG agar plates, the closest relative in known species, the maximum similarity with the closest relative, its classification in the Clostridiales cluster, origin of mouse ID, and culture medium for isolation.

Table 3 shows, for each of 31 bacterial strains isolated from the caecal contents of mouse #F8, #G2, #H1 and #K3 using BL agar or EG agar plates, the closest relative in known species, the maximum similarity with the closest relative, the database used for BLAST search, and similarity between strains.

Table 4 shows 16S rDNA analysis for each of 31 strains that were isolated. Bacterial DNA was isolated from each of the 31 strains and the 16S rDNA of the isolates was amplified by colony-PCR. Each amplified DNA was purified, sequenced, and aligned using the ClustalW software program. Based on the sequence of 16S rDNA for each strain, their closest species, % similarity with the closest species, and the similarity to other strains are shown. Strains that were included in the 23-mix, 17-mix, P-mix, 5 mix-B, 5-mix-C, and 3 mix are marked in the right hand column.

DETAILED DESCRIPTION

<Composition Having Effect of Inducing Proliferation or Accumulation of Regulatory T Cells>

Described herein is a composition that induces proliferation, accumulation of regulatory T cells or both proliferation and accumulation of regulatory T cells. The composition comprises, as an active ingredient, one or more of the following: a (at least one, one or more) organism selected from the group consisting of: Clostridium saccharothermum, Clostridium ramosum JCM1298, Clostridium ramosum, Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_5FAA, Lachnospiraceae bacterium 6_1_63FAA, Clostridium sp. 14616, Clostridium bolteae ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes cococeae, Clostridium bolteae, Lachnospiraceae bacterium DJF_VP30, Lachnospiraceae bacterium 3_1_57FAA_CT1, Anaerotruncus colihominis, Anaerotruncus colihominis DSM 17241, Ruminococcus sp. IDB, Lachnospiraceae bacterium 2_1_46FAA, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum WAL-14163, Eubacterium ontorum, Clostridium sp. D5, Oscillibacter bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 31600208, and Clostridiales bacterium 1_7_47FAA, Blautia cocoides, Anaerostipes cac-
cae DSM 14662, a culture supernatant of one or more of the bacteria, a component of culture medium in which a (at least one, one or more) bacterium described herein has grown, a physiologically active substance derived from a (at least one; one or more) bacterium described herein; and a (at least one, one or more) bacterium containing DNA comprising a nucleotide sequence having at least 97% homology to the nucleotide sequence of DNA of any of the bacterial species described herein, such as those listed above. Bacteria described herein were isolated from human fecal samples using the methods outlined in Examples 19 to 28.

The term “regulatory T cells” refers to T cells that suppress an abnormal or excessive immune response and play a role in immune tolerance. The regulatory T cells are typically transcription factor Foxp3-positive CD4-positive T cells. The regulatory T cells of the present invention also include transcription factor Foxp3-negative regulatory T cells that are IL-10-producing CD4-positive T cells.

The term “induces proliferation or accumulation of regulatory T cells” refers to an effect of inducing the differentiation of immature T cells into regulatory T cells, which differentiation leads to the proliferation and/or the accumulation of regulatory T cells. Further, the meaning of “induces proliferation or accumulation of regulatory T cells” includes in-vivo effects, in vitro effects, and ex vivo effects. All of the following effects are included: an effect of inducing in vivo proliferation or accumulation of regulatory T cells through administration or ingestion of the aforementioned bacteria belonging to the Clostridia class, a culture supernatant of the bacteria or supernatant component(s), or a physiologically active substance derived from the bacteria; an effect of inducing proliferation or accumulation of cultured regulatory T cells by causing the aforementioned bacteria belonging to the Clostridia class, a culture supernatant of the bacteria or supernatant component(s), or a physiologically active substance derived from the bacteria to act on the cultured regulatory T cells; and an effect of inducing proliferation or accumulation of regulatory T cells which are collected from a living organism and which are intended to be subsequently introduced into a living organism, such as the organism from which they were obtained or another organism, by causing the aforementioned bacteria belonging to the Clostridia class, a culture supernatant of the bacteria or supernatant component(s), or the physiologically active substance derived from the bacteria to act on the regulatory T cells. The effect of inducing proliferation or accumulation of regulatory T cells can be evaluated, for example, as follows. Specifically, the aforementioned bacteria belonging to the Clostridia class, a culture supernatant of the bacteria or supernatant component(s), or a physiologically active substance derived from the bacteria is orally administered to an experimental animal, such as a germ-free mouse, and CD4-positive cells in the colon are isolated, and the ratio of regulatory T cells contained in the CD4-positive cells is measured by flow cytometry (refer to Example 7).

The regulatory T cells whose proliferation or accumulation is induced by the composition of the present invention are preferably transcription factor Foxp3-positive regulatory T cells or IL-10-producing regulatory T cells.

In the present invention, “human-derived bacteria” means bacterial species that have been isolated from a fecal sample or from a gastrointestinal biopsy obtained from a human individual or whose ancestors were isolated from a fecal sample or from a gastrointestinal biopsy obtained from a human (e.g., are progeny of bacteria obtained from a fecal sample or a gastrointestinal biopsy). For example, the bacterial species may have been previously isolated from a fecal sample or from a gastrointestinal biopsy obtained from a human and cultured for a sufficient time to generate progeny. The progeny can then be further cultured or frozen. The human-derived bacteria are naturally occurring commensals that populate the gastrointestinal tract of human individuals, preferably healthy human individuals.

In the present invention, the term “Clostridia class” (as in “compositions containing bacteria belonging to the Clostridia class”) refers to a class of Gram-positive, obligate anaerobic bacteria belonging to the Firmicutes phylum that have the ability to form spores. It is important to note that while currently most bacteria in this class are included in the Clostridiales order, this categorization is still partly based on old methods and is likely to be redefined in the future based on new advances in sequencing technologies that are enabling sequencing of the full genomes of bacteria in this class. Table 2 provides a summary of the categorization of 17 abundant species belonging to the Clostridia class which have been identified by the inventors as strong Treg-inducers and cultured in vitro. All of these species fall, under current classification rules, in the Clostridiaceae family, and belong to clusters IV, XIVA, XVIA, and XVIII.

The composition of the present invention may include one strain alone (only one strain) of any of the aforementioned bacterial species, but two or more strains of the bacteria can be used together. For example, one, two, three, four, five, six, seven, eight, nine, ten, eleven, twelve, thirteen, fourteen, fifteen, sixteen or seventeen of the strains listed in Table 2 or Table 4, in any combination, can be used together to affect regulatory T cells. In some embodiments, the 23, 17, 5, or 3 species mixes listed in Table 4 can be used together (and administered in one or several compositions) to affect regulatory T cells. In some embodiments, the following strains can be combined (the composition comprises): strain 1 (OTU136, closest species: Clostridium saccharogranum, Clostridium ramosum JCM1298), strain 2 (OTU221, closest species: flavinreductase plantii, Pseudoflavinreductase capillosus ATCC 29799), strain 4 (OTU9, closest species: Clostridium hathewayi, Clostridium saccharolyticum WM1), strain 5 (OTU296, closest species: Clostridium scindens, Lachnospiraceae bacterium 5.1-57F(AA)), strain 6 (OTU21, closest species: Blautia cocoides, Lachnospiraceae bacterium 6.1-63F(AA)), strain 7 (OUT 166, closest species: Clostridium sp., Clostridium bolteae ATCC BAA-613), strain 8 (OTU73, closest species: cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2.2.44A), strain 9 (OTU174, closest species: Clostridium indolis, Anaerostipes caccae DSM 14662), strain 10 (OTU166, closest species: Clostridium bolteae, Clostridium bolteae ATCC BAA-613), strain 12 (OTU55, closest species: Lachnospiraceae bacterium DJF VP30, Lachnospiraceae bacterium 3.1-57F(AA)_CT1), strain 13 (OTU337, closest species: Anaerotruncus colihominis, Anaerotruncus colihominis DSM 17241), strain 14 (OTU314, closest species: Ruminococcus sp. ID8, Lachnospiraceae bacterium 2.1-46F(AA)), strain 15 (OTU195, closest species: Clostridium lavalense, Clostridium asparagiforme DSM 15981), strain 16 (OTU306, closest species: Clostridium symbiosum, Clostridium symbiosum WAL-14163), strain 18 (OTU46, closest species: Clostridium ramosum, Clostridium ramosum), strain 21 (OTU87, closest species: Eubacterium contourtorum, Clostridium sp. D5), strain 23 (OTU152, closest species: Lachnospiraceae bacterium DJF VP30, Lachnospiraceae bacterium 3.1-57F(AA)_CT1), strain 24 (OTU255, closest species: Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes), strain 25 (OTU259, closest species: Eubac-
terium contortum, Clostridium sp. D5), strain 26 (OTU281, closest species: Clostridium scindens, Lachnospiraceae bacterium 5_1_57F(A)A), strain 27 (OTU288, closest species: Lachnospiraceae bacterium A4, Lachnospiraceae bacterium 3_1_57F(A)A_CT1), strain 28 (OTU344, closest species: Clostridiales bacterium 1_7_47F(A)A), and strain 29 (OTU359, closest species: Lachnospiraceae bacterium A4, Lachnospiraceae bacterium 3_1_57F(A)A_CT1) as described in Table 4.

In some embodiments, the following strains can be combined (the composition comprises): strain 1 (OTU136, closest species: Clostridium saccharogenum, Clostridium ramosum JCM1298), strain 3 (OTU221, closest species: Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799), strain 4 (OTU9, closest species: Clostridium hathewayi, Clostridium saccharolyticum WM1), strain 6 (OTU21, closest species: Blautia cocoides, Lachnospiraceae bacterium 6_1_63F(A)A), strain 7 (OUT 166, closest species: Clostridium sp., Clostridium botulinum ATCC BAA-613), strain 9 (OTU174, closest species: Clostridium indolis, Anaerostipes caccae DSM 15981), strain 28 (OTU344, closest species: Clostridium sp., Clostridium botulinum ATCC 29799) as described in Table 4. In some embodiments, the following strains can be combined: strain 1 (OTU136, closest species: Clostridium saccharogenum, Clostridium ramosum JCM1298), strain 2 (OTU46, closest species: Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799) and strain 3 (OTU221, closest species: Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799) as described in Table 4. The use of multiple strains of the aforementioned species of bacteria, preferably belonging to the Clostridiaceae cluster XIVa, the cluster IV in combination can bring about an excellent effect on regulatory T cells. In addition to the bacteria belonging to clusters XIVa and IV, Clostridium ramosum, Clostridium saccharogenum (belonging to cluster XVIII) and cf. Clostridium sp., MLG055 (belonging to cluster XVIII) can also be used. If more than one strain of bacteria is used (e.g., one or more strain belonging to cluster XIVa, one or more strain belonging to cluster IV, one or more strain belonging to clusters XVIII or XVI or a combination of any of the foregoing), the number and ratio of strains used can vary widely. The number and ratio to be used can be determined based on a variety of factors (e.g., the desired effect, such as induction or inhibition of proliferation or accumulation of regulatory T cells, the disease or condition to be treated, prevented or reduced in severity; the age or gender of the recipient; the typical amounts of the strains in healthy humans). The strains can be present in a single composition, in which case they can be consumed or ingested together (in a single composition), or can be present in more than one composition (e.g., each can be in a separate composition), in which case they can be consumed individually or the compositions can be combined and the resulting combination (combined compositions) consumed or ingested. Any number or combination of the strains that proves effective (e.g., any number from one to 22, such as 1 to 20, 1 to 15, 1 to 10, 1 to 5, 1 to 3, 1 to 2, and any number therebetweentwo or one to 23, such as 1 to 23, 3 to 23, 5 to 23, 1 to 20, 1 to 17, 3 to 17, 5 to 17, 1 to 15, 1 to 10, 1 to 5, 3 to 5, 1 to 3, 1 to 2, and any number therebetweentwo) can be administered. In certain embodiments of the present invention, a combination of some or all of the 22 or 23 (e.g., the 23 strains in Example 32 and Table 4) strains described in the present disclosure is used. For example, at least one, two or more, three, three or more, four, four or more, five, five or more, six, six or more or any other number of the 22 or 23 described strains, including 22 or 23 strains, can be used. In some embodiments, the specific combinations of 3, 5, 17, or 23 strains described in Table 4 can be used (the composition comprises combinations of 3, 5, 17 or 23 strains described in Table 4). They can be used in combination with one another and in combination with strains not described in the cited reference.

Cells of bacteria belonging to the Clostridial class, such as these specifically described herein, can be used in spore form or in vegetative form. From the viewpoint of stability to high temperature and pressure conditions, extended shelf life, ease of handling, resistance to antibiotics, and lack of need for a cold chain storage and distribution, the bacteria may be preferably in the form of spore. From the viewpoint of abiding by the directives of certain manufacturing organizations that do not tolerate spore contamination in their
facilities, the bacteria may alternatively be produced (and later administered) in the form of vegetative cells.

The term “physiologically active substance derived from bacteria belonging to the Clostridia class” of the present invention includes substances contained in the bacteria, secretion products of the bacteria, and metabolites of the bacteria. Such a physiologically active substance can be identified by purifying an active component from the bacteria, a culture supernatant thereof, or intestinal tract contents in the intestinal tract of a mouse in which only bacteria belonging to the Clostridia class are colonized by an already known purification method.

“Chloroform treatment” of a fecal sample obtained from a human is a method that isolates the bacteria in the fecal sample that have the ability to form spores, and is not particularly limited, as long as the spore-forming fraction is obtained by treating feces of a human with chloroform (for example, 3% chloroform), and has the effect of inducing proliferation or accumulation of regulatory T cells, including mammalian regulatory T cells such as murine regulatory T cells and human regulatory T cells.

When the aforementioned “bacteria belonging to the Clostridia class” are cultured in a medium, substances contained in the bacteria, secretion products and metabolites produced by the bacteria are released from the bacteria. The meaning of the active ingredient “culture supernatant of the bacteria” in the composition of the present invention includes such substances, secretion products, and metabolites. The culture supernatant is not particularly limited, as long as the culture supernatant has the effect of inducing proliferation or accumulation of regulatory T cells. Examples of the culture supernatant include a protein fraction of the culture supernatant, a polysaccharide fraction of the culture supernatant, a lipid fraction of the culture supernatant, and a low-molecular weight metabolite fraction of the culture supernatant.

The bacterial composition may be administered in the form of a pharmaceutical composition, a dietary supplement, or a food or beverage (which may also be an animal feed), or may be used as a reagent for an animal model experiment. The pharmaceutical composition, the dietary supplement, the food or beverage, and the reagent induce proliferation or accumulation of regulatory T cells. An example presented herein revealed that regulatory T cells (Treg cells) induced by bacteria or the like belonging to the Clostridia class suppressed the proliferation of effector T-cells. The composition of the present invention can be used suitably as a composition having an immunosuppressive effect. The immunosuppressive effect can be evaluated, for example, as follows. Regulatory T cells isolated from an experimental animal, such as a mouse, to which the composition of the present invention is orally administered are caused to act on effector T-cells (CD4+ CD25+ cells) isolated from the spleen, and the proliferation ability thereof is measured by using the intake amount of [3H]-thymidine as an index (refer to Example 14).

The bacterial composition of the present invention can be used, for example, as a pharmaceutical composition for preventing or treating (reducing, partially or completely, the adverse effects of) an autoimmune disease, such as chronic inflammatory bowel disease, systemic lupus erythematosus, rheumatoid arthritis, multiple sclerosis, or Hashimoto’s disease; an allergic disease, such as a food allergy, pollenosis, or asthma; an infectious disease, such as an infection with Clostridium difficile; an inflammatory disease such as a TNF-mediated inflammatory disease (e.g., an inflammatory disease of the gastrointestinal tract, such as pouchitis, a cardiovascular inflammatory condition, such as atherosclerosis, or an inflammatory lung disease, such as chronic obstructive pulmonary disease); a pharmaceutical composition for suppressing rejection in organ transplantation or other situations in which tissue rejection might occur; a supplement, food, or beverage for improving immune functions; or a reagent for suppressing the proliferation or function of effector T-cells.

More specific examples of target diseases for which the composition is useful for treatment (reducing adverse effects or prevention) include autoimmune diseases, allergic diseases, infectious diseases, and rejection in organ transplantations, such as inflammatory bowel disease (IBD), ulcerative colitis, Crohn’s disease, sprue, autoimmune arthritis, rheumatoid arthritis, Type I diabetes, multiple sclerosis, graft vs. host disease following bone marrow transplantation, osteoarthritis, juvenile chronic arthritis, Lyme arthritis, psoriatic arthritis, reactive arthritis, spondyloarthropathy, systemic lupus erythematosus, insulin dependent diabetes mellitus, thyroiditis, asthma, psoriasis, dermatitis, dermatis, graft versus host disease, acute or chronic immune disease associated with organ transplantation, sarcoidosis, atherosclerosis, disseminated intravascular coagulation, Kawasaki’s disease, Grave’s disease, nephrotic syndrome, chronic fatigue syndrome, Wegener’s granulomatosis, Henoch-Schoenlein purpura, microscopic vasculitis of the kidneys, chronic active hepatitis, uveitis, septic shock, toxic shock syndrome, sepsis syndrome, cachexia, acquired immunodeficiency syndrome, acute transverse myelitis, Huntington’s chorea, Parkinson’s disease, Alzheimer’s disease, stroke, primary biliary cirrhosis, hemolytic anemia, polyglanular deficiency type I syndrome and polyglanular deficiency type II syndrome, Schmidt’s syndrome, adult (acute) respiratory distress syndrome, allopuria, alopecia areata, seronegative arthropathy, arthropathy, Reiter’s disease, psoriatic arthropathy, chlamydia, versinis and salmonella associated arthropathy, spondyloarthropathy, atheromatous disease/arteriosclerosis, allergic colitis, atopic allergy, food allergies such as peanut allergy, tree nut allergy, egg allergy, milk allergy, soy allergy, wheat allergy, seafood allergy, shellfish allergy, or sesame seed allergy, autoimmune bulous disease, pemphigus vulgaris, pemphigus foliaceus, pemphigoid, linear IgA disease, autoimmune haemolytic anaemia, Coombs positive haemolytic anaemia, acquired pernicious anaemia, juvenile pernicious anaemia, myalgic encephalitis/Royal Free Disease, chronic mucocutaneous candidiasis, giant cell arteritis, primary sclerosing hepatitis, cryopgenic autoimmune hepatitis, Acquired Immunodeficiency Disease Syndrome, Acquired Immunodeficiency Related Diseases, Hepatitis C, common varied immunodeficiency (common variable hypogammaglobulinaemia), dilated cardiomyopathy, fibrotic lung disease, cryptogenic fibrosing alveolitis, postinflammatory interstitial lung disease, interstitial pneumonitis, connective tissue disease associated interstitial lung disease, mixed connective tissue disease associated lung disease, systemic sclerosis associated interstitial lung disease, rheumatoid arthritis associated interstitial lung disease, systemic lupus erythematosus associated lung disease, dermatomyositis/polymyositis associated lung disease, Sjogren’s disease associated lung disease, ankylosing spondyloarthritis associated lung disease, vasculitic diffuse lung disease, haemosideros associated lung disease, drug-induced interstitial lung disease, radiation fibrosis, bronchiolitis obliterans, chronic eosinophilic pneumonia, lymphocytic infiltrative lung disease, postinfectious interstitial lung disease, gouty arthritis, autoimmune hepatitis, type-I autoimmune hepatitis (classical autoimmune or...
lupoid hepatitis), type-2 autoimmune hepatitis (anti-LKM antibody hepatitis), autoimmune mediated hypoglycemia, type B insulin resistance with acanthosis nigricans, hyperparathyroidism, acute immune disease associated with organ transplantation, chronic immune disease associated with organ transplantation, osteoarthrosis, primary sclerosing cholangitis, idiopathic lupus, autoimmune neuroptenia, renal disease NOS, gomelurenoneritides, microscopic vasulitis of the kidneys, discoid lupus, erythematous, male infertility idiopathic or NOS, sperm autoimmunity, multiple sclerosis (all subtypes), insulin-dependent diabetes mellitus, sympathetic ophthalmia, pulmonary hypertension secondary to connective tissue disease, Goodpasture’s syndrome, pulmonary manifestation of polymyositis nodosa, acute rheumatoid fever, rheumatoid spondylitis, Still’s disease, systemic sclerosis, Takayasu’s disease/arteritis, autoimmune thrombocytopenia, idiopathic thrombocytopenia, autoimmune thyroid disease, hyperthyroidism, goitrous autoimmune hypothyroidism (Hashimoto’s disease), atrophic autoimmune hypothyroidism, primary myxoedema, phacomatous uveitis, primary vasculitis, vitiligo, allergic rhinitis (pollen allergies), anaphylaxis, pet allergies, latex allergies, drug allergies, allergic rhinoconjunctivitis, eosinophilic esophagitis, hypereosinophilic syndrome, eosinophilic gastroenteritis cutaneous lupus erythematosus, eosinophilic esophagitis, hypereosinophilic syndrome, and eosinophilic gastroenteritis, and diarrhea.

Additional examples of target diseases for which the composition is useful for treatment include colon cancer, cystic fibrosis, celiac disease, Type 2 diabetes, and autism-related immunopathologies. These diseases are characterized by a reduction of Clostridium Clusters IV and XIV in the gastrointestinal microbiota.

Compositions described herein can also be used as a pharmaceutical composition for preventing or treating infectious diseases in an individual whose resistance to the infectious diseases is impaired, for example because of damage due to excessive inflammation caused by the immunity or due to an alteration of the patient’s microbiome. Examples of infectious pathogens that impair maintenance or recovery of homeostasis of a host, and which eventually bring about such immunopathological tissue damage include Salmonella, Shigella, Clostridium difficile, Mycobacterium (which cause the disease tuberculosis), protozoa (which cause malaria), filarial nematodes (which cause the disease filariasis), Schistosoma (which cause schistosomiasis), Toxoplasma (which cause the disease toxoplasmosis), Leishmania (which cause the disease leishmaniasis), HCV and HBV (which cause the disease hepatitis C and hepatitis B), and herpes simplex viruses (which cause the disease herpes).

Pharmaceutical preparations can be formulated from the bacterial compositions described by drug formulation methods known to those of skill in the art. For example, the composition can be used orally or parenterally in the form of capsules, tablets, pills, sachets, liquids, powders, granules, fine granules, film-coated preparations, pellets, troches, sublingual preparations, chewables, buccal preparations, pastes, syrups, suspensions, elixirs, emulsions, liniments, ointments, plasters, cataplasms, transdermal absorption systems, lotions, inhalations, aerosols, injections, suppositories, and the like.

For formulating these preparations, the bacterial compositions can be used in appropriate combination with carriers that are pharmacologically acceptable or acceptable for ingestion, such as in a food or beverage, including one or more of the following: sterile water, physiological saline, vegetable oil, solvent, a base material, an emulsifier, a suspending agent, a surfactant, a stabilizer, a flavoring agent, an aromatic, an excipient, a vehicle, a preservative, a binder, a diluent, a toxicity adjusting agent, a soothing agent, a bulking agent, a disintegrating agent, a buffer agent, a coating agent, a lubricant, a colorant, a sweetener, a thickening agent, a flavor corrigent, a solubilizer, and other additives.

A pharmaceutical preparation or formulation and particularly a pharmaceutical preparation for oral administration, comprises an additional component that enables efficient delivery of the bacterial composition of the present invention to the colon, in order to more efficiently induce proliferation or accumulation of regulatory T cells in the colon. A variety of pharmaceutical preparations that enable the delivery of the bacterial composition to the colon can be used. Examples thereof include pH sensitive compositions, more specifically, buffered solution formulations or enteric polymers that release their contents when the pH becomes alkaline after the enteric polymers pass through the stomach. When a pH sensitive composition is used for formulating the pharmaceutical preparation, the pH sensitive composition is preferably a polymer whose pH threshold of the decomposition of the composition is between about 6.8 and about 7.5. Such a numeric value range is a range in which the pH shifts toward the alkaline side at a distal portion of the stomach, and hence is a suitable range for use in the delivery to the colon.

Another embodiment of a pharmaceutical preparation useful for delivery of the bacterial composition to the colon is one that ensures the delivery to the colon by delaying the release of the contents (e.g., the bacterial composition) by approximately 3 to 5 hours, which corresponds to the small intestinal transit time. In one embodiment of a pharmaceutical preparation for delayed release, a hydrogel is used as a shell. The hydrogel is hydrated and swells upon contact with gastrointestinal fluid, with the result that the contents are effectively released (released predominantly in the colon). Delayed release dosage units include drug-containing compositions having a material which coats or selectively coats a drug or active ingredient to be administered. Examples of such a selective coating material include in vivo degradable polymers, gradually hydrolyzable polymers, gradually water-soluble polymers, and/or enzyme degradable polymers. A wide variety of coating materials for efficiently delaying the release is available and includes, for example, cellulose-based polymers such as hydroxypropyl cellulose, acrylic acid polymers and copolymers such as methacrylic acid polymers and copolymers, and vinyl polymers and copolymers such as polyvinylpyrrolidone.

Examples of the composition enabling the delivery to the colon further include bioadhesive compositions which specifically adhere to the colonic mucosal membrane (for example, a polymer described in the specification of U.S. Pat. No. 6,368,586) and compositions into which a protease inhibitor is incorporated for protecting particularly a biopharmaceutical preparation in the gastrointestinal tracts from decomposition due to an activity of a protease.

An example of a system enabling the delivery to the colon is a system of delivering a composition to the colon by pressure change in such a way that the contents are released by utilizing pressure change caused by generation of gas in bacterial fermentation at a distal portion of the stomach. Such a system is not particularly limited, and a more specific example thereof is a capsule which has contents dispersed in a suppository base and which is coated with a hydrophobic polymer (for example, ethyl cellulose).
Another example of the system enabling the delivery to the colon is a system of delivering a composition to the colon, the system being specifically decomposed by an enzyme (for example, a carbohydrate hydrolase or a carbohydrate reductase) present in the colon. Such a system is not particularly limited, and more specific examples thereof include systems which use food components such as non-starch polysaccharides, amylose, xanthan gum, and azopolymers.

When used as a pharmaceutical preparation, the bacterial composition may be used in combination with an already known pharmaceutical composition for use in immunosuppression. In some embodiments, the pharmaceutical preparation can comprise both the bacterial composition and the already known pharmaceutical composition. Such a known pharmaceutical composition is not particularly limited, and may be at least one therapeutic composition selected from the group consisting of corticosteroids, mesalazine, mesalamine, sulfasalazine, sulfasalazine derivatives, immunosuppressive drugs, cyclosporin A, mercaptopurine, azathioprine, prednisone, methotrexate, anti-histamines, glucocorticoids, epinephrine, theophylline, Cromolyn sodium, anti-leukotrienes, anti-cholinergic drugs for rhinitis, anti-cholinergic decongestants, mast-cell stabilizers, monoclonal anti-IgE: antibodies, vaccines (preferably vaccines used for vaccination where the amount of an allergen is gradually increased), anti-TNF inhibitors such as infliximab, adalimumab, certolizumab pegol, golimumab, or etanercept, and combinations thereof. It is preferable to use these therapeutic compositions in combination with the bacterial composition described herein. The bacterial composition can also be used as an adjuvant to improve the efficacy of a vaccine formulation such as a vaccine for the prophylaxis or treatment of an autoimmune disease or an allergic disease.

The bacterial composition can be used as a food or beverage, such as a health food or beverage, a food or beverage for infants, a food or beverage for pregnant women, athletes, senior citizens or other specified group, a functional food, a beverage, a food or beverage for specified health use, a dietary supplement, a food or beverage for patients, or an animal feed. Specific examples of the foods and beverages include various beverages such as juices, refreshing beverages, tea beverages, drink preparations, jelly beverages, and functional beverages; alcoholic beverages such as beers; carbohydrate-containing foods such as rice food products, noodles, breads, and pastas; paste products such as fish hams, sausages, paste products of seafood; retort pouch products such as curries, food dressed with a thick starchy sauces, and Chinese soups; soups; dairy products such as milk, dairy beverages, ice creams, cheeses, and yogurts; fermented products such as fermented soybean pastes, yogurts, fermented beverages, and pickles; bean products; various confectionary products such as Western confectionary products including biscuits, cookies, and the like; Japanese confectionary products including steamed bean-jam buns, soft adzuki-bean jellies, and the like, candies, chewing gums, marmalade, cold desserts including jellies, crème caramels, and frozen desserts; instant foods such as instant soups and instant soy-bean soups; microwavable foods; and the like. Further, the examples also include health foods and beverages prepared in the forms of powders, granules, tablets, capsules, liquids, pastes, and jellies. The composition of the present invention can be used for animals, including humans. The animals, other than humans, are not particularly limited, and the composition can be used for various livestock, poultry, pets, experimental animals, and the like. Specific examples of the animals include pigs, cattle, horses, sheep, goats, chickens, wild ducks, ostriches, domestic ducks, dogs, cats, rabbits, hamsters, mice, rats, monkeys, and the like, but the animals are not limited thereto.

Without wishing to be bound by theory, individuals in whom bacteria belonging to the group Firmicutes (the group to which the Clostridium clusters IV and VII belong) are relatively abundant gain more body weight than individuals in whom bacteria belonging to the group Bacteroidetes are relatively abundant. The bacterial composition is capable of conditioning absorption of nutrients and improving feed efficiency. From such a viewpoint, the bacterial composition can be used for promoting body weight gain, or for a high efficiency animal feed. Diseases and conditions that would benefit from body weight gain include, e.g., starvation, cancer, AIDS, gastrointestinal disorders (e.g., celiac disease, peptic ulcer, inflammatory bowel disease (Crohn’s disease and ulcerative colitis), pancreatitis, gastritis, diarrheaea), hyperthyroidism, infection, renal disease, cardiac disease, pulmonary disease, connective tissue disease, weight loss caused by medications, anorexia, Addison’s disease, dementia, depression, hypercalcemia, Parkinson’s disease and tuberculosis.

The addition of the bacterial composition to an antibiotic-free animal feed makes it possible to increase the body weight of an animal that ingests the animal feed to a level equal to or higher than that achieved by animal ingesting antibiotic-containing animal feeds, and also makes it possible to reduce pathogenic bacteria in the stomach to a level equal to those in animals consuming typical antibiotic-containing animal feeds. The bacterial composition can be used as a component of an animal feed that does not need the addition of antibiotics.

In addition, unlike conventional bacteria (Lactobacillus and Bifidobacteria) in commercial use, which are not easy to incorporate into the livestock production, the present bacterial composition in spore form can be pelleted, sprayed, or easily mixed with an animal feed and can also be added to drinking water.

Animal feed comprising the bacterial composition can be fed to a wide variety of types of animals and animals of a varying ages and can be fed at regular intervals or for a certain period (for example, at birth, during weaning, or when the animal is relocated or shipped).

The bacterial composition can be used to promote weight gain and enhance energy absorption in humans and nonhumans (e.g., farm or other food animals).

The bacterial active components of the bacterial composition can be manufactured using fermentation techniques well known in the art. In one embodiment, the active ingredients are manufactured using anaerobic fermentors, which can support the rapid growth of bacterial species belonging to the Clostridia class. The anaerobic fermentors may be, for example, stirred tank reactors or disposable wave bioreactors. Culture media such as BL, media and EG media, or similar versions of these media devoid of animal components can be used to support the growth of the bacterial species. The bacterial product can be purified and concentrated from the fermentation broth by traditional techniques, such as centrifugation and filtration, and can optionally be dried and lyophilized by techniques well known in the art.

A food or beverage comprising a bacterial composition described herein can be manufactured by manufacturing techniques well known in the technical field. One or more components (for example, a nutrient) which are effective for the improvement of an immune function by an immuno-sup-
pressive effect may be added to the food or beverage. In addition, the food or beverage may be combined with another component or another functional food exhibiting a function other than the function of the improvement of an immune function to thereby serve as a multi-functional food or beverage.

Moreover, the bacterial composition can be incorporated into foods requiring a processing step which may destroy ordinary probiotic strains. Specifically, the most commercially usable probiotic strains cannot be incorporated into foods that need to be processed, for example, by heat treatment, long term storage, freezing, mechanical stress, or high-pressure treatment (for example, extrusion forming or roll forming). On the other hand, because of the advantageous nature of forming spores, the bacterial composition described herein can be easily incorporated into such processed foods. For example, the bacterial composition in the form of spores can survive even in a dried food, and can remain living even after being ingested. The bacterial composition can withstand low-temperature sterilization processes; typically processes carried out at a temperature from about 70°C to about 100°C, both inclusive. The bacterial composition can be incorporated into dairy products that require a pasteurization step. Furthermore, the bacterial composition can withstand long-term storage of many years; high-temperature processing such as baking and boiling; low-temperature processing such as freezing and cold storage; and high-pressure treatments such as extrusion forming and roll forming.

Many types of foods that need to be processed under such harsh conditions include foods which need to be processed in a microwave oven to be edible (for example, oatmeal), foods which need to be baked to be edible (for example, a muffin), foods which need to be subjected to a sterilization high-temperature treatment for a short period of time to be edible (for example, milk), and foods which need to be heated to be drinkable (for example, hot tea).

The amount of the bacterial composition to be administered or ingested can be determined empirically, taking into consideration such factors as the age, body weight, gender, symptoms, health conditions, of an individual who will receive it, as well as the kind of bacterial composition (a pharmaceutical product, a food or beverage) to be administered or ingested. For example, the amount per administration or ingestion is generally 0.01 mg/kg body weight to 100 mg/kg body weight, and, in specific embodiments, 1 mg/kg body weight to 10 mg/kg body weight. Also described herein is a method for suppressing the immunity (reducing the immune response) of a subject, the method being characterized in that the bacteria belonging to the Clostridia class or the physiologically active substance derived from the bacteria is administered to or ingested by the subject as described above.

The bacterial composition may be administered to an individual once, or it may be administered more than once. If the composition is administered more than once, it can be administered on a regular basis (for example, once a day, once every two days, once a week, once every two weeks, once a month, once every 6 months, or once a year) or on an as needed or irregular basis. The appropriate frequency of administration (which may depend on host genetics, age, gender, and health or disease status of the subject, among other factors) may be determined empirically. For example, a patient can be administered one dose of the composition, and the levels of the bacterial strains of the composition in fecal samples obtained from the patient can be measured at different times (for example after 1 day, after 2 days, after 1 week, after 2 weeks, after 1 month). When the levels of the bacteria fall to, for example, one half of their maximum value, a second dose can be administered, and so on.

A product comprising the bacterial composition (a pharmaceutical product, a food or beverage, or a reagent) or a manual thereof may be accompanied by a document or statement explaining that the product can be used to suppress the immunity (including a statement that the product has an immunosuppressive effect and a statement that the product has an effect of suppressing the proliferation or function of effector T cells). Here, the “provision to the product or the manual thereof with the note” means that the document or statement is provided to a main body, a container, a package, or the like of the product, or the note is provided to a manual, a package insert, a leaflet, or other printed matters, which disclose information on the product.

(Method for Inducing Proliferation or Accumulation of Regulatory T Cells)

As described above, and as shown in Examples, administration of the bacterial composition to an individual makes it possible to induce proliferation or accumulation of regulatory T cells in the individual. This provides a method of inducing proliferation or accumulation of regulatory T cells in an individual, the method comprising: administering, to the individual, at least one substance selected from the group consisting of: (a) Clostridium saccharogena, Clostridium ramosum JCM1298, Clostridium ramosum, Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1.57F5AA, Lachnospiraceae bacterium 6_1.63F6AA, Clostridium sp. 14616, Clostridium bolteae ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2.44A, Clostridium indolis, Anaerostipes caccae, Clostridium bolteae, Lachnospiraceae bacterium D9F_VP30, Lachnospiraceae bacterium 3.1.57F6AA_C1T, Anaerotruncus colihominis, Anaerotruncus colihominis DSM 1724, Ruminococcus sp. IDB, Lachnospiraceae bacterium 2_1.46F6AA, Clostridium lavalense, Clostridium asporagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum W14-14163, Eubacterium centorum, Clostridium sp. D5, Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002D8, and Clostridiales bacterium 1.7.47F6AA, Blautia cocoides, Anaerostipes caccae DSM 14662; (b) a culture supernatant of at least one (a, one or more) of the bacteria described/listed herein; (c) a physiologically active substance derived from a (one or more, at least one) bacterium described/listed herein; or a combination of any two or three of (a), (b) and (c). The bacterial composition is administered (provided) to the individual in sufficient quantity to produce the desired effect of inducing proliferation, accumulation or both proliferation and accumulation of regulatory T cells. It may be administered to an individual in need of treatment, reduction in the severity of or prevention of at least one disease selected from an autoimmune disease, an inflammatory disease, an allergic disease, and an infectious disease.

Note that, the “individual” or “subject” may be in a healthy state or a diseased state. The method may further comprise the optional step of administering at least one (a, one or more) antibiotic preceding, or in combination with, the bacterial composition. The antibiotic administered can be, for example, one which facilitates recolonization of the gut by Gram-positive bacteria of the Clostridia class, such as an antibiotic that reduces Gram-negative bacteria. Examples

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of such antibiotics include aminoglycoside antibiotics (aminoglycoside antibiotics (ami-
kanamycin, neomycin, netilmicin, tobramycin, and paromomycin), cephalosporin antibiotics
(cefalexin, cefadroxil, cefetan, cefprozil, cefuroxime, cefixime, cefditoren, cefepime, cefotaxime,
ceftriaxone, cefotetan, ceftriaxone, cefotaxime, cefuroxime, and cefotaxime), sulfonamides, ampicillin, and streptomycin).

Moreover, a probiotic composition such as almond skin, inulin, oligofructose, raffinose, lactulose, pectin, hemicellulose (such as xyloglucan and alpha-glucans), amylopectin, and resistant starch which are not decomposed in the upper gastrointestinal tract and promote the growth of intestinal microbes in the intestinal tract, as well as growth factors such as acetyl-Co A, biotin, beet molasses, and yeast extracts, preferentially contributes to the proliferation of the bacterial species in the composition belonging to the Clostridial class. A method of inducing proliferation and/or accumulation of regulatory T cells in an individual can comprise administering, to the individual, at least one substance selected from the above in combination with the bacterial composition. Also contemplated herein is a composition comprising the bacterial composition and a probiotic composition.

The above-described antibiotic, and the above-described probiotic composition or growth factor may be used in combination. Moreover, a therapeutic composition may be administered to an individual together with at least one substance selected from the group consisting of the bacterial composition, an antibiotic, and a probiotic composition or growth factor.

A therapeutic composition can be, for example, one therapeutic composition selected from the group consisting of corticosteroids, mesalazine, mesalamine, sulfasalazine, sulphasalazine derivatives, immunosuppressive drugs, cyclosporin A, mercuric pentamine, azathioprine, prednisone, methotrexate, antithymine, glucocorticoids, epinephrine, threophylline, cromolyn sodium, anti-cho-
linergic drugs for rhinitis, anti-cholinergic decongestants, mast-cell stabilizers, monoclonal anti-lg antibodies, vaccines (preferably, vaccines used for vaccination where the amount of an allergen is gradually increased), anti-TNF inhibitors such as infliximab, adalimumab, certolizumab pegol, golimumab, or etanercept, and combinations thereof. These therapeutic compositions can be administered prior to, in combination with or following administration of the bacterial composition and optionally, also in combination with an antibiotic, a probiotic composition, a growth factor or any combination of an antibiotic, a probiotic composition and a growth factor.

There is no particular limitation imposed on the combined use of the therapeutic composition with at least one substance selected from the group consisting of the bacterial composition, the “antibiotic”, and the “probiotic composition or growth factor”. For example, the “one substance” and the therapeutic composition are administered orally or parenterally to an individual simultaneously or sequentially/ individually at any appropriate time.

Whether administration of the bacterial composition induces the proliferation and/or accumulation of regulatory T cells can be determined by using, as an index, increase or reinforcement of at least one of the following: the number of regulatory T cells, the ratio of regulatory T cells in the T cell group of the colon, a function of regulatory T cells, or expression of a marker of regulatory T cells. A specific approach is measurement counts or percentage of Foxp3-expressing Tregs in a patient sample, such as a biopsy or a blood sample, promotion (enhancement) of IL-10 expres-
sion, promotion (enhancement) of CTLA4 expression, promotion (enhancement) of IDO expression, suppression of IL-4 expression, or colonization of an individual with the bacterial composition administered as the index of the induction of proliferation or accumulation of regulatory T cells.

Methods for detecting such expression include northern blotting, RT-PCR, and dot blotting for detection of gene expression at the transcription level; ELISA, radioimmuno-
assays, immunoblotting, immunoprecipitation, and flow cytometry for detection of gene expression at the translation level.

Samples that may be used for measuring such an index include tissues and fluids obtained from an individual, such as blood, obtained in a biopsy, and a fecal sample.

Method for Predicting Response of an Individual to the Bacterial Composition by Monitoring the Individual’s Response to Treatment with the Composition

Also described is a method in which an amount (e.g. count) or the percentage of at least one bacterial species selected from the group consisting of: Clostridium sphenoca-
orugum, Clostridium ramosum, Flavonifractor plautii, Pseudoflavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57F7A, Lachnospiraceae bacterium 6_1_63F8A, Clostridium sp. 14616, Clostridium botalec ATCC BAA-613, ef, Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium ino-
lis, Anaerotipes caccae, Clostridium botalec, Lachnospiraceae bacterium DIF_VP3, Lachnospiraceae bacteri-
unum 3_1_57F7A, CT1, Anaerotruncus colitহiminis, Anaerotruncus colitহiminis DSM 17241, Ruminococcus sp. IDB, Lachnospiraceae bacterium 2_1_46F8A, Clostridium lavalense, Clostridium asparagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum Wal-
14163, Eubacterium conter, Clostridium sp. D5, Oscil-
ospiraceae bacterium NML 061048, Oscillibacter valeri-
genesis, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47F8A, Blaxi-
tia acidol i, Anaerotipes caccae DSM 14662 in a patient’s sample (e.g. a colonic biopsy or a fecal sample) is determined. When the percentage or the count of the bacteria selected from the list above is lower in an individual than a base line value obtained by performing a similar determination on a healthy individual (e.g., an individual who does not have/has not been identified as having a disease or condition for which the bacterial composition is a potential treatment such as an autoimmune disease, an allergic condition, cancer, organ rejection), it is determined that the individual is likely to be responsive to the bacterial composition. This determination can be used, for example, by a clinician to determine whether an individual or a patient is likely to benefit from treatment with the bacterial composition, or to select an individual or a patient for inclusion in a clinical trial. The clinician can then administer the bacterial composition to the individual or patient based on the determination that the individual or patient is likely to benefit from treatment. This determination can also be used as a method to monitor an individual’s response to treatment with the bacterial compositions described, wherein a higher value of the determination after treatment with the bacterial composition (compared to a determination before treatment) indicates that the individual has responded favorably to treatment (e.g. a positive indicator of successful colonization and enhanced immunosuppression in the individual).
 Optionally, the prognosis and monitoring methods described here may further comprise the step of measuring in the individual's samples the percentages or absolute counts of other commensal species belonging to Clostridium Clusters IV and XIVa that are not present in the bacterial composition, wherein lower than baseline values before treatment indicate a higher likelihood of a positive response to treatment, and wherein an increased value after treatment indicates that the individual has responded favorably to treatment. In the prognosis and monitoring methods described here, a variety of known methods can be used for determining the composition of the microbiota. For example, 16S rRNA sequencing can be used.

“Vaccine Adjuvant Composition and Method for Treating or Preventing Infectious Disease or Autoimmune Disease by Using the Vaccine Composition”

As described above, and as shown in the Examples, the induction of Treg cells in the colon by bacteria belonging to the Clostridia class has an important role in local and systemic immune responses. The bacterial composition can also be used as an adjuvant to improve the efficacy of a vaccine formulation. In one embodiment, the bacterial composition can be used as an adjuvant to a vaccine for the prophylaxis or treatment of an autoimmune disease or an allergic disease (for example, as an adjuvant for a vaccination protocol where the amount of an allergen is gradually increased).

Example of autoimmune diseases and allergic diseases include those described as the “specific examples of target diseases” in “Composition Having Effect of Inducing Proliferation or Accumulation of Regulatory T cells”.

Other Embodiment

The bacterial composition can also be administered to an individual who is also receiving antibiotic treatment. The present inventors have demonstrated that antibiotics that act against Gram+ bacteria, such as vancomycin or metronidazole, can effectively eliminate or greatly reduce bacterial species belonging to the Clostridia class from the gastrointestinal tract of mammals and subsequently decrease the levels of regulatory T cells (Example 5, FIG. 30). Without wishing to be bound by theory, the key role of bacteria belonging to the Clostridia class in preserving immune tolerance strongly indicates that their absence or reduced levels can play a key role in autoimmune diseases characterized by failures of immune tolerance. Accordingly, individuals undergoing courses of antibiotics against Gram+ bacteria (for example, individuals being treated for infections with pathogens such as C. difficile and Giardia), who are at a high risk of experiencing a loss of the bacteria belonging to the Clostridia class and thus experience immune tolerance deficits, can be prevented by “repopulating” through use of the bacterial composition. The bacterial composition can be administered before, simultaneously with, or after the antibiotic treatment, but preferably it is administered simultaneously or after the antibiotic treatment. The bacterial composition is preferably administered in spore form, to improve its resistance to residual antibiotics. Antibiotics against Gram-positive bacteria include, but are not limited to, vancomycin, metronidazole, linezolid, ramoplanin, fidaxomicin, cephalosporin antibiotics (cephalexin, cefuroxime, cefadroxil, cefazolin, cephalothin, cefaclor, cefamandole, cefotin, cefprozil, and cefotibiprole); fluoroquinolone antibiotics (cipro, Levaquin, floxin, tequin, avelox, and norflox); tetracycline antibiotics (tetracycline, minocycline, oxytetracycline, and doxycycline); penicillin antibiotics (amoxicillin, ampicillin, penicillin V, dicloxacillin, carbenicillin, vancomycin, and methicillin); and carbapenem antibiotics (ertapenem, doripenem, imipenem/ cilastatin, and meropenem).

“Methods to Select Treg-Inducing Organisms”

Also described is a method of obtaining bacteria capable of inducing Tregs, comprising: (1) isolating the bacterial spore-forming fraction from a fecal or biopsy sample obtained from a mammal, preferably a human (e.g. by chloroform treatment or by heat treatment), (2) optionally, orally administering the spore-forming fraction to a non-human mammal, preferably a germ-free non-human mammal; (3) optionally, obtaining a fecal sample from the non-human mammal, diluting the fecal sample (for example diluting it by volume by a factor of 10, 100, 1,000, or 10,000), thereby producing a diluted fecal sample, and orally administering the diluted sample to a second germ-free non-human mammal, wherein optional step (3) can be repeated more than one time, (4) plating serial dilutions, under aerobic condition or strictly anaerobic conditions, of either the spore-forming fraction obtained in (1) or a sample of intestinal contents of the non-human mammal of (3), and (5) picking a single colony from the culture plate. The colony can be further assessed for the ability of bacteria to induce proliferation of regulatory T cells and/or accumulation of regulatory T cells using known methods, such as those described in the examples.

Following are examples, which describe specific aspects. They are not intended to be limiting in any way.

Note that mice used in Examples were prepared or produced as follows. In the following description, mice may be referred to as “SPI” or “GF”. These “SPI” and “GF” indicate that the mice were maintained in the absence of specific pathogenic bacteria (specific pathogen-free, SPF), and that the mice were maintained under Germ-Free (GF) conditions, respectively.

“Mice”

C57BL/6, Balb/c, and I/J mice maintained under SPF or GF conditions were purchased from Sankyo Labo Service Corporation, Inc. (Japan), JAPAN SLC, INC. (Japan), CLEA Japan, Inc. (Japan), or The Jackson Laboratory (USA). GF mice and gnotobiotic mice were bred and maintained within the gnotobiotic facility of The University of Tokyo, YakuI Central Institute for Microbiological Research, or Sankyo Labo Service Corporation, Inc. Myd88−/−, Rip2−/−, and Card9−/− mice were produced as described in NPL 1 to 3, and backcrossed for 8 generations or more, so that a C57BL/6 genetic background was achieved. Foxp3−/− mice were purchased from the Jackson Laboratory.

“II10<sup>trans</sup> Mice”

To form a bicistronic locus encoding both II10 and Venus under control of an II10 promoter, a targeting construct was first created. Specifically, a cassette (ires-Venus-SV40 polyA signal cassette, refer to Non-Patent Document 4) which was made of an internal ribosome entry site (ires), a yellow fluorescent protein (Venus), and a SV40 polyA signal (SV40 polyA) and which was arranged next to a neomycin-resistant gene (neo), was inserted between a stop codon and a polyA signal (Eoxon 5) of a II10 gene. Next, the obtained targeting construct was used to cause homologous recombination with the II10 gene region in the genome of mice. Thus, II10<sup>trans</sup> mice having an II10<sup>trans</sup> allele were produced (refer to FIG. 1). Note that in FIG. 1, “<sup>trans</sup>” represents a gene coding thymidine kinase, “<sup>neo</sup>” represents the neomycin-resistant gene, and “<sup>BamH1</sup>” represents a cleavage site by the restriction enzyme BamH1.
Genomic DNAs were extracted from the Il10<sup>−/−</sup> mice, treated with BamHI, and Southern blotted by use of a probe shown in FIG. 1. FIG. 2 shows the obtained results. Wild-type and Il10<sup>−/−</sup> alleles were detected as bands having sizes of 19 kb and 5.5 kb, respectively. Hence, as is apparent from the results obtained, the homozygous recombination occurred in the genome of the Il10<sup>−/−</sup> mice.

Further, CD4<sup>+</sup> Venus<sup>+</sup> cells or CD4<sup>+</sup> Venus<sup>+</sup> cells in the colonic lamina propria of the Il10<sup>−/−</sup> mice were sorted by use of a FACSAria. Then, real-time RT-PCR was carried out on an ABI 7300 system by a method to be described later, to determine the amount of IL-10 mRNA expressed. It was found that, since the development of the IL-10 mRNA was detected only in the CD4<sup>+</sup> Venus<sup>+</sup> cells, the expression of IL-10 mRNA in the Il10<sup>−/−</sup> mice was correctly reflected in the expression of Venus. Note that the germ-free states of such Il10<sup>−/−</sup> mice were established in Central Institute for Experimental Animals (Kawasaki, Japan). The Il10<sup>−/−</sup> mice in the germ-free states were maintained in vinyl isolators in Sankyo Labo Service Corporation, Inc. (Tokyo, Japan), and used in the following Examples.

Experiments and analyses in Examples were carried out as follows.

<Method for Colonization of Mice with Murine Bacteria and Analysis Thereof>

According to the description in NPL 5 and 6, mice in which SFB or Clostridium were colonized were produced. Cecal contents or feces of the obtained gnotobiotic mice were dissolved in sterile water or an anaerobic dilution solution. The dissolved cecal contents or feces as they were or after a chloroform treatment were orally administered to GF mice. Three strains of the Lactobacillus and 16 strains of the Bacteroides were cultured separately from each other in a BL or EGl agar medium in an anaerobic manner. The cultured bacteria were harvested, suspended in an anaerobic TS broth, and orally administrated forcibly to GF mice. The state of the colonization of the bacteria in the mice was assessed by microscopic observation conducted on a smear preparation of fecal pellets.

<Isolation of Intestinal Lamina Propria Lymphocytes and Flow Cytometry>

The small intestine and colon were collected and opened longitudinally. The cecum was also isolated and cecal content was directly frozen at −80°C or suspended in 2 ml PBS, then added 40% glycerol (final concentration 20%), snap-frozen in liquid nitrogen and stored at −80°C until use. The colon and small intestine were washed in PBS to remove all luminal contents and shaken in Hank’s balanced salt solution (HBSS) containing 5 mM EDTA for 20 min at 37°C. After removing epithelial cells, muscle layers and fat tissue using tweezers, the lamina propria layers were cut into small pieces and incubated with RPMI1640 containing 4% fetal bovine serum, 1 mg/ml collagenase D, 0.5 mg/ml dispase and 40 μg/ml DNase I (all Roche Diagnostics) for 1 h at 37°C in a shaking water bath. The digested tissues were washed with HBSS containing 5 mM EDTA, resuspended in 5 ml of 40% Percoll (GE Healthcare) and overlaid on 2.5 ml of 80% Percoll in a 15-ml Falcon tube. Percoll gradient separation was performed by centrifugation at 800 g for 20 min at 25°C. The lamina propria lymphocytes were collected from the interface of Percoll gradient and suspended in ice-cold PBS. For analysis of regulatory T cells, isolated lymphocytes were labeled with the LIVE/DEAD fixable violet dead cell stain kit (Invitrogen) to exclude dead cells in the analysis. The cells were washed with staining buffer containing PBS, 2% FBS, 2 mM EDTA and 0.09% NaN3 and stained surface CD4 with PE/Cy7-labeled anti-CD4 Ab (RM4-5, BD Biosciences). Intracellular staining of Foxp3 and Helios was performed using the Alexa700-labeled anti-Foxp3 Ab (FJK-16s, eBioscience), Alexa647-labeled anti-Helios (22F6, eBioscience) and Foxp3 Staining Buffer Set (eBioscience). For analysis of Th1 and Th17 cells, isolated lymphocytes were stimulated for 4 hours with 50 ng/ml phorbol 12-myristate 13-acetate (PMA, Sigma) and 1 μg/ml ionomycin (Sigma) in the presence of GolgiStop (BD Biosciences). After incubation for 4 hours, cells were washed in PBS, labeled with the LIVE/DEAD fixable violet dead cell stain kit and stained surface CD4 with PE/Cy7-labeled anti-CD4 Ab. Cells were washed, fixed in Cytotox/Cytoperm, permeabilized with Perm/Wash buffer (BD Biosciences), and stained with the APC-labeled anti-IL-17 Ab (eBio1B7, eBioscience) and FITC-labeled anti-IFN-γ Ab (XMG1.2, BD Biosciences). The Ab stained cells were analyzed using a LSR Fortessa (BD Biosciences), and data were analyzed using FlowJo software (TreeStar).

<Real-Time RT-PCR>

From an RNA prepared by using RNAeasy Mini Kit (Qiagen), a cDNA was synthesized by use of a MMV reverse transcriptase (Promega KK). The cDNA obtained was analyzed by real-time RT-PCR using Power SYBR Green PCR Master Mix (Applied Biosystems) and ABI 7300 real-time PCR system (Applied Biosystems), or real-time RT-PCR using SYBR Premix Ex Taq (TAKARA) and Light Cycler 480. For each sample, a value obtained was normalized for the amount of GAPDH. A primer set was designed by using Primer Express Version 3.0 (Applied Biosystems), and those exhibiting a 90% or higher sequence identity at an initial evaluation were selected. The primer set used was as follows:

**Fopx3**

5′-GCGACATAGTTCTCCCGAGATT-3′ *(SEQ ID NO: 1)*

5′-GGCTGCAATATTGTGTACTTG-3′ *(SEQ ID NO: 2)*

**CTLA4**

5′-CCCTTTTGATGCCTGTACCT-3′ *(SEQ ID NO: 3)*

5′-GGCTACCCGTATGCTCCAG-3′ *(SEQ ID NO: 4)*

**GTR**

5′-TCAATGGCAAAGCTGCAAGCA-3′ *(SEQ ID NO: 5)*

5′-ACACCGGAAACCCAACACA-3′ *(SEQ ID NO: 6)*

**IL-10**

5′-GATTCTGAAGTGTCAGGAGT-3′ *(SEQ ID NO: 7)*

5′-CTCTTATGAGTGGTTAGGATGTCGA-3′ *(SEQ ID NO: 8)*

**GAPDH**

5′-CCCTCGTCCCCGTAGCAAAAA-3′ *(SEQ ID NO: 9)*

5′-TTCCTCACCTGGCAGTCAAC-3′ *(SEQ ID NO: 10)*

**Mmp2**

5′-GGCAATGTCTTTCTGAGGAC-3′ *(SEQ ID NO: 11)*
<Preparation and Culturing of Large Intestinal Epithelial Cells (IECs)>  
First, the colon was collected, cut open longitudinally, and rinsed with PBS. Subsequently, the colon was treated with 1 mM dithiothreitol (DTT) at 37°C for 30 minutes on a shaker, and then vortexed for one minute to disrupt the epithelial integrity. The released intestinal epithelial cells (IECs) were collected, and suspended in 5 ml of 20% percoll. The suspension was overlayed on 2.5 ml of 80% percoll in a 15-ml Falcon tube. Then, the tube was centrifuged at 25°C and 780 g for 20 minutes to conduct cell separation by percoll density gradient centrifugation. Cells at the interface were collected, and used as colonic IECs (purity: 90% or higher, viability: 95%). The IECs obtained were collected and suspended in RPMI containing 10% FBS, and 1x10⁶ cells of the IECs were cultured in a 24-well plate for 24 hours. Then, the culture supernatant was collected, and measured for active TGF-β1 level by ELISA (Promega).

Meanwhile, for culturing T cells in vitro, 1.5x10⁵ MACS-purified splenic CD4⁺ T cells were cultured in each well of a round-bottomed 96-well plate, together with a 50% conditioned medium in which IECs isolated from GF mice or Clostridium-colonized mice were cultured, and with 25 ng/ml of hIL-2 (Peprotech), in the presence or absence of 25 μg/ml of an anti-TGF-β antibody (R&D). Note that 10 μg/ml of an anti-CD3 antibody and an anti-CD28 antibody (BD Bioscience) were bound to the round-bottomed plate. After a 5-day culture, the CD4⁺ T cells were collected, and subjected to a real-time PCR.

<Colitis Experimental Model>
A fecal suspension from Clostridium-colonized mice was orally administered to C57BL/6 mice (2-week old), which were grown in a conventional environment for six weeks.

For preparing a DSS-induced colitis model, 2% (wt/vol) DSS (reagent grade, DSS salt, molecular weight=36 to 50 kDa, manufactured by MP Biomedicals), together with drinking water, was given to the mice for six days.

Meanwhile, for preparing an oxazolone-induced colitis model, the mice were pre-sensitized by transdermally applying, onto the mice, 150 μl of a 3% oxazolone (4-ethoxyethylene-2-phenyl-2-oxazolin-5-one, Sigma-Aldrich)/100% ethanol solution. Five days after that, 150 μl of a 1% oxazolone/50% ethanol solution was intrarectally administered again to the pre-sensitized mice under a light anesthesia. Note that the intrarectal administration was conducted by using a 3.5 F catheter.

Each mouse was analyzed daily for body weight, occult blood, bleeding visible with the naked eyes (gross blood), and the hardness of stool. Moreover, the body weight loss percentage, intestinal bleeding (no bleeding, occult blood (hemocult+), or bleeding visible with the naked eyes), and the hardness of stool (normal stool, loose stool, or diarrhea) were evaluated numerically, and the disease activity index (DAI) was calculated in accordance with the description in “S. Wirtz, C. Neufert, B. Weigmann, M. F. Neurath, Nat Protoc 2, 541 (2007).”

<OVA-Specific IgE Reaction>
BALB/c SPF mice were inoculated with a fecal suspension from Clostridium-colonized mice (2-week old), and grown in a conventional environment. Then, 1 μg of OVA (grade V, Sigma) and 2 mg of alum (Thermo Scientific), 0.2 ml in total, were intraperitoneally injected to the mice (at their ages of 4 weeks and 6 weeks). Sera were collected every week from the mice at the root of their tail, and OVA-specific IgE was measured by ELISA (Chondrex). Then, at their ages of 8 weeks, splenic cells were collected, inoculated in a 96-well plate at 1x10⁶ cells per well, and stimulated with OVA (100 μg/ml) for three days. Thereafter, the culture supernatant was collected, and measured for IL-4 and IL-10 levels by ELISA (R&D).

<Statistical Analysis>
The difference between control and experimental groups was evaluated by the Student’s t-test.

<Chloroform Treatment and Oral Inoculation with Fecal Samples into GF Mice>
Human stool (2 g) from a healthy volunteer (Japanese, male, 29 y old) was suspended with 20 ml phosphate-buffered saline (PBS) and passed through a 70 μm cell strainer to eliminate clumps and debris. Then fecal suspension was mixed with or without chloroform (final concentration 3%), and incubated in a shaking water bath for 60 min. The fecal suspensions without chloroform treatment were orally inoculated into germ-free (GF) mice (250 μl/mouse). After evaporation of chloroform by bubbling with N2 gas for 30 min, the aliquots containing chloroform-resistant (spore-forming) fraction of human intestinal bacteria were inoculated into IQI GF mice. Each group of ex-GF mice was separately kept in a vinyl isolator for 3 or 4 weeks.

<Co-Housing Experiment>
To evaluate whether Treg-inducing human bacteria can be transmitted horizontally, IQI GF mice were co-housed for 4 weeks with ex-GF mice colonized with chloroform-treated human feces (Example 21 mice) in a vinyl isolator (6 mice, designated as mouse #D1 to #D6).

Inoculation with Diluted Cecal Contents Into GF Mice>
The frozen cecal content from ex-GF mice inoculated with chloroform-treated human feces (#C4) was suspended in 10 times volume (w/v) of PBS, passed through a 70 μm cell strainer and treated 3% chloroform. Then the suspension was diluted 2000 (for 4 mice, designated as mouse #E1 to #E4) or 20000 (for 8 mice, designated as mouse #F1 to #F8) by PBS and orally inoculated into GF IQI mice (2.5x10⁸ or 2.5x10⁹ cells/250 μl/mouse). After 4 weeks, lymphocytes were collected from colon and small intestine and analyzed for Foxp3+ Treg cell proportion and their Helios expression. Cecal contents were frozen and stored at -80°C until use.
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<Re-Colonization Experiments>

The frozen cecal content from ex-GF mice inoculated with 20000-fold dilution (#F3, 7 and 8) was suspended in 10 times volume (w/v) of PBS, passed through a 70 μm cell strainer and treated 3% chloroform. The suspensions were

orally inoculated into GF IqI mice (5, 4 or 4 mice; designated as mouse #G1 to #G5, #H1 to #H4 or #I1 to #I4, respectively). After 4 weeks, colon and small intestine were collected and analyzed for Foxp3+ Treg cell proportion and their Helios expression. Cecal contents were suspended in 20% glycerol solution, snap-frozen in liquid nitrogen and stored at −80°C.

<Cultured Bacteria-Colonization Experiments>

The glycerol stock of cecal content from #G2 mouse was diluted with PBS and seeded onto BL agar plate. After 48 hours, all bacterial colonies were collected by scraping the plates with a plate scraper and inoculated into GF IqI mice (4 mice, designated as mouse #K1 to #K4). Six bacterial strains were isolated from the freeze stock of cecal content from #F8 mouse using BL agar plate. These isolated strains were inoculated into GF IqI mice (4 mice, designated as mouse #J1 to #J4). (Details of the culture method are described below.)

<16S rRNA Gene Quantitative PCR Analysis>

Using a QiAamp DNA Stool mini kit (QiAGEN), bacterial genomic DNA was isolated from the human stool from a healthy volunteer as described above (human stool), cecal contents from GF mice gavaged with chloroform-treated human stool (cecal content of B4 mouse) or feces from SPF ICR mouse (feces of SPF mouse). The isolated DNA was used as template for quantitative PCR. The amplification program consisted of one cycle at 95°C for 1 min, followed by 50 cycles at 95°C for 10 s and 60°C for 30 s. Quantitative PCR analysis was carried out using a Light-Cycler 480 (Roche). Relative quantity was calculated by the ΔΔCt method and normalized to the amount of total bacteria. The following primer sets were used: total bacteria, 5'-GGTG-

GAAATACGTCCCGG-3' (SEQ ID NO.: 45) and 5'-TACGG-

CTACCCGTTTTAGACCT-3' (SEQ ID NO.: 46); Clostridium cluster XIVa (Clostridium cocoides subgroup), 5'-AAAAGCCCCTGGATGATCA-3' (SEQ ID NO.: 47) and 5'-CTTGTGAGTTTCACTTCGTGCA-3' (SEQ ID NO.: 48); Clostridium cluster XIV b (Clostridium leptum) 5'-CCCT-

TGGCGGAGGAGTCC-3' (SEQ ID NO.: 49) and 5'-AAGATTTACCCACTTACTGGCCT-3' (SEQ ID NO.: 50); Bacteroides, 5'-GGAGGAGGAAGGTCCACCC-3' (SEQ ID NO.: 51) and 5'-GCTTACCTGGCTGTTCCAG-3' (SEQ ID NO.: 52); Bifidobacterium, 5'-CGGTTGAG-

TAATGCGGTTGAC-3' (SEQ ID NO.: 53) and 5'-TGATAGGACCCGACCAC-3' (SEQ ID NO.: 54). Note that mice gavaged with chloroform-treated human stool exhibited high levels of spore-forming bacteria, such as Clostridium clusters XIVa and IV, and a severe decrease of non-spore-forming bacteria, such as Bacteroides and Bifo-

dobacterium, compared with the human stool before chloro-

form treatment.

<Isolation of DNA from Cecal Contents for 16S RNA Gene Metasquence Analysis>

The cecal contents of A1-1, A2-4, B-4, E-3, E-7, E-8, F-2, G-3, H-3, I-3 and J-3 were collected by centrifugation at 5000g for 10 min at 4°C, suspended in 10 ml of Tris-

EDTA containing 10 mM Tris-HCl and 1 mM EDTA (pH 8), and then used for DNA isolation. Lysozyme (SIGMA, 15 mg/ml) was added to the cell suspension. After incubation at 37°C for 1 h with gentle mixing, a purified achromopeptidase (Wako) was added (final 2000 units/ml) and incubated at 37°C for 30 min. Then, sodium dodecyl sulfate (final

1%) was added to the cell suspension and mixed well. Subsequently, proteinase K (Merck) was added (final 1 mg/ml) to the suspension and the mixture was incubated at 55°C for 1 h.

High-molecular-weight DNA was isolated and purified by phenol/chloroform extraction, ethanol, and finally polyethy-

lene glycol precipitation.

<16S rRNA Gene Metasquence>

An aliquot of the DNA was used for PCR amplification and sequencing of bacterial 16S rDNA genes. ~330 bp amplicons, spanning variable region 1-2 (V1-2) of the gene were generated by using (i) modified primer 8F (5'-CCATCTCCATCCCTGCGTGTTCGACTAGCAG-Bac-

code+aggtgttagtmtgtagcag-3' (SEQ ID NO.: 55)) which consists of 454 adaptor sequence (underlined), a sample specific, error correcting barcode (10 bases, bold) and the universal bacterial primer 8F and (ii) modified primer 338R (5'-CTATATCCCCGTGGTACCTAGGAGATGTG-3' (SEQ ID NO.: 56)) which contains 454 adaptor sequence (underlined) and the bacterial primer 338R. Polymerase chain reactions were performed for each focal DNA sample: each 0.5-μl reaction contained 40 ng of DNA, 5 μl of 10× Ex Taq buffer (TAKARA), 5 μl of 2.5 mM dNTP mixture, 0.2 μl Ex Taq and 0.2 μM of each primer. PCR conditions consisted of an initial denaturation step performed at 96°C for 2 min, followed by 20 cycles of denaturation (96°C, 30 s), annealing (55°C, 30 s) and amplification (72°C, 1 min) and final amplification step performed at 72°C for 10 min. Amplicons generated from each sample were subsequently purified using AMPure XP (Beckman Coulter). The amount of DNA was quantified using Quant-IT Picogreen dsDNA Assay Kit (Invitrogen) and TBS-380 mini Fluorometer (Turner Bioskysm). The amplified DNA were used as template for 454 GS Junior (Roche) pyrosequencing. The sequences were performed using GS Junior Titanium emPCR Kit-Lib1-L, GS Junior Titanium Sequencing Kit and GS Junior Titanium PicoTi-

ter Plate Kit (all Roche) according to the manufacturer's manuals (GS Junior Titanium Series, emPCR Amplification Method Manual—Lib1-L and Sequencing Method Manual). Resulting sequences (3400 reads were produced for each sample) were classified into OTU on the basis of sequence similarity (>97% identity). Representative sequences from each OTU were compared with sequences in nucleic acid databases (Ribosomal Database Project) using BLAST to determine the closest relatives. Then, OTUs were classified into species on the basis of the closest relatives. All data of close relatives and the number of reads are shown in Table 1.

1. <Isolation of Bacterial Strains>

Bacterial strains isolated from the cecal contents of #F8, #G2, #H1 and #K3 by plating serial dilutions of the cecal samples under aerobic condition or strictly anaerobic condition (80% N2 10% H2 10% CO2) onto BL agar (Eiken Chemical) or EG agar plates containing medium with the following components (quantities expressed per liter): Meat extract 500 ml; Proteose peptone No. 3 (10.0 g, Difco); Yeast Extract (5.0 g, Difco); Na2HPO4 (4.0 g); D(-)+Glucose (1.5 g); Soluble Starch (0.5 g); L-cysteine (0.5 g); L-cysteine-HCl—H2O (0.5 g); Tween80 (0.5 g); Bacto Agar (16.0 g, Difco); defibrinated horse blood (50 ml). After culture at 37°C for 2 or 4 days, each single colony was picked up and cultured for additional 2 or 4 days at 37°C by ABCM broth or EG agar plate. The isolated strains were collected into EG stock medium (10% DMSO) and stored at −80°C. For suspension of isolated strains to re-inoculate mice, TS medium (27.5 g of trypticase soy broth w/o
dextrose. 0.84 g of Na2CO3, 0.5 g of L-cysteine-HCl—
H2O, 1000 ml of distilled water, pH adjusted to 7.2+/-0.2
with NaOH; then autoclaved for 15 minutes at 115 degrees
Celsius. To identify the isolated strains, 16S rRNA coding
sequence genes were performed. The 16SrRNA genes were
amplified by colony PCR using KOD FX (TOYOBO), 16S
rRNA gene-specific primer pairs: 5’-AGGTGTTAGTCC-
MTGCTAGCAG-3’ (SEQ ID NO.: 57) and 519R (5’-ATAC-
CCGCGTCTG-3’ (SEQ ID NO.: 58)) for C. indolis, C.
boletae, Bacteroides sp. MANG, L. bacterium D1F-VSP30,
A. clostibomis, Ruminococcus sp. IDB, C. lavalense, C.
symbiosis and E. coli. contortum or 1513R (5’-ACGGCTAC-
CTTGTACGACTTT-3’ (SEQ ID NO.: 59)) for C. saccha-
rogumia, C. ramosum, F. plautii, C. halleyaei, C. scindens,
Clostridium sp. 2335, Clostridium sp. 14616 and of
Clostridium sp. MLG055 and GeneAmp PCR System9700
(Appplied Biosystems). The amplification program consisted
of one cycle at 98°C, for 2 min, followed by 40 cycles at 98°
C. for 10 s, 56°C, for 30s and 68°C, for 40 s. Each amplified
DNA was purified from the reaction mixture using Illustra
GFX PCR DNA and Gel Band Purification Kit (GE Health-
care). Sequence analysis was performed using BigDye Ter-
minator V3.1 Cycle Sequencing Kit (Applied Biosystems)
and Applied Biosystems 3730xl DNA analyzer (Applied
Biosystems). The resulting sequences were compared with
sequences in nucleic acid databases using BLAST to deter-
mine the closest relatives. The closest relatives and %
identity of all isolated strains, information for genus-species
of the closest relatives, Clostridium cluster, ID of mouse
from which was derived, maximum similarity and culture
medium of isolated strains were summarized in Table 2.

EXAMPLE 1

First, it was investigated whether or not accumulation of
regulatory T cells (Treg cells) in the colonic lamina propria
was dependent on commensal bacteria. Specifically, lympho-
cytes were isolated from peripheral lymph nodes (pLN) of
Balb/c mice bred in the absence of specific pathogenic
bacteria (SPF) or from lamina propria of the colon or the
small intestine (SI) of the mice. The CD4 and Foxp3 were
stained by antibodies. Then, the ratio of Foxp3+ cells in
CD4+ lymphocytes was analyzed by flow cytometry. The
results showed that Foxp3+ Treg cells were present at a high
frequency in the lamina propria of the gastrointestinal tract,
especially in the colonic lamina propria, of the mice kept
under the environment free from specific pathogenic micro-
organisms (SPF). In addition, it was also found that the
number of the Foxp3+ Treg cells in the colonic lamina
propria gradually increased up to three months after
birth, whereas the number of the Foxp3+ Treg cells in the
periodic lymph nodes was basically constant from the time
of two weeks after their birth.

EXAMPLE 2

Next, it was investigated whether or not the temporal
accumulation of the Treg cells in the colon in found in
Example 1 had a relationship with the colonization of
intestinal commensal microbiota. Specifically, the expres-
sion of CD4 and the expression of Foxp3 in lymphocytes
isolated from the small intestine, the colon, and the periph-
eral lymph nodes of mice bred under a germ-free (GF) or
SPF environment (8 weeks old: Balb/c mice, IQL mice, and
C57BL/6 mice) were analyzed. Similar results were
obtained in three or more independent experiments.

In addition, lamina propria lymphocytes were collected
from SPF mice and Gf1 mice (Balb/c mice or C57BL/6 mice).
CD4 and Foxp3 were stained with antibodies. Then, the
lamina propria lymphocytes were analyzed by FACS.

Further, lymphocytes were isolated from the lamina prop-
ria of the colon, the lamina propria of the small intestine
(SI), Peyer’s patches (PPs), and mesenteric lymph nodes
(MLNs) of mice (SPF C57BL/6 mice) to which antibiotics
were orally administered with water for eight weeks. CD4
and Foxp3 were stained with antibodies. Then, the lym-
phocytes were analyzed by FACS. Similar results (the ratio
of the Foxp3+ cells in the CD4+ cells of an individual mouse)
were obtained in two or more independent experiments.
Note that the following antibiotics were used in combination
in accordance with the description in the following docu-
ment:
ampicillin (A; 500 mg/L, Sigma)
vancomycin (V; 500 mg/L, NACALAI TESQUE, INC.)
metronidazole (M; 1 g/L, NACALAI TESQUE, INC.)
neomycin (N; 1 g/L, NACALAI TESQUE, INC.)
Rakoff-Nahoum, J. Paglino, F. Eslami-Varzaneh, S. Edberg,
R. Medzhitov, Cell 118, 229 (Jul. 23, 2004)
Fagarasan et al., Science 298, 1424 (Nov. 15, 2002)
As is apparent from the results, the frequencies and the
absolute numbers of Foxp3+ CD4+ cells in the small intesti-
ne and the peripheral lymph nodes of the GF mice were
equal to or greater than those of the SPF mice. In addition,
the numbers of the Treg cells in the small intestinal lamina
propria, Peyer’s patches, and mesenteric lymph nodes of
the SPF mice to which the antibodies were orally adminis-
tered for eight weeks were equal to or greater than those of
the SPF mice that had not received antibiotics. Meanwhile,
the number of the Foxp3+ CD4+ cells in the colonic lamina
propria of the GF mice was decreased significantly in
comparison with that of the SPF mice. This decrease was
commonly observed among mice of different genetic back-
grounds (Balb/c, IQL, and C57BL/6), as well as among mice
bred in different animal facilities. In addition, it was also
shown that the number of Treg cells in the colonic lamina
propria of the SPF C57BL/6 mice to which the antibodies
were administered was decreased significantly.

EXAMPLE 3

Next, it was directly checked whether or not the decrease
in the number of the Treg cells in the colonic lamina propria
of the GF mice shown in Example 2 was attributed to the
absence of microbiota. Specifically, a fecal suspension of B6
SPF mice purchased from The Jackson Laboratory was
orally administered to GF-IQL mice (conventionalization).
Three weeks after the administration, lymphocytes were
isolated from the colonic lamina propria, and the expres-
sion of Foxp3 in CD4+ lymphocytes was analyzed. The results
showed that the number of Treg cells in the small intestinal
lamina propria did not change. However, the number of the
Treg cells in the colonic lamina propria increased signifi-
cantly. Hence, it was shown that host-microbial interaction
played an important role in the accumulation of Foxp3+ Treg
cells in the colonic lamina propria, while the accumulation
of the Treg cells in the small intestinal lamina propria had a
different mechanism.

EXAMPLE 4

Next, the relationship between the gut-associated lymph-
oid tissues of mice and the number of Foxp3+ cells in the
colonic lamina propria of the mice was investigated in
accordance with the method described in M. N. Kweon et al., J Immunol 174, 4565 (Apr. 1, 2005). Specifically, 100 µg
of an extracellular domain recombinant protein (a fusion protein (LTβR-lg) between a lymphotoxin β receptor
(LTβR) and a Fc region of human IgG1, refer to Honda et al., J Exp Med 193, 621 (Mar. 5, 2001)) was injected
intraperitoneally into pregnant C57Bl/6j mice 14 days after conception. The LTβR-lg was again injected intraperitone-
ally into fetuses obtained from such mice, so that mice from which isolated lymphoid follicles (ILFs), Peyer’s patches
(PPs), and colonic-patches (CPs) were completely removed were produced. Then, the ratios of Foxp3+ cells in CD4+ cells
in the colonic lamina propria of the mice treated with the LTβR-lg, and mice treated with rat IgG (control) were
analyzed by FACS. The results show that the ratio of the Foxp3+ cells in the colonic lamina propria of the mice
deficient in isolated lymphoid follicles, Peyer’s patches, and the colonic-patches (the mice treated with the LTβR-lg)
rather increased. Accordingly, it was suggested that the decrease in the number of Treg cells in the colonic
lamina propria of the GF mice and the mice treated with the antibiotics was caused because the transmission of specific
signals which promotes the accumulation of Treg cells in the colonic lamina propria and which is caused by the intestinal
microbes did not occur, rather than simply because of a secondary effect of disorganized gut-associated lymphoid
tissues.

EXAMPLE 5

To investigate whether or not a specific intestinal flora induced the accumulation of colonic Treg cells, vancomycin
as an antibiotic against Gram-positive bacteria or polymyxin B as an antibiotic against Gram-negative bacteria was
administered to SPF mice (from 4 weeks of age) for four weeks, and analyzed for the ratio of Foxp3+ cells in
the CD4+ cell group (% Foxp3+ in CD4).

The results show that the number of Treg cells in the colon of the mice to which vancomycin was administered was
markedly decreased in comparison with that of the control. In contrast, no influence was observed on the number of
Treg cells of the mice to which polymyxin B was admin-
istered. Those facts suggested that Gram-positive commensal bacteria played a major role in accumulation of Treg cells.

EXAMPLE 6

A recent report has suggested that spore-forming bacteria play an important role in intestinal T cells response (see V.
Gaboriau-Routhiau et al., Immunity 31, 677 (Oct. 16, 2009)). In this respect, fecal microorganisms (spore-forming
fraction) resistant to 3% chloroform were orally admin-
istered to GF mice, which were then analyzed for the ratio of Foxp3+ cells in the CD4+ cell group (% Foxp3+ in CD4).

Three weeks after the administration of the chloroform-
treated feces, the number of Treg cells in the administered
mice was markedly increased to the same level as those of the
SPF mice and the GF mice to which the untreated feces
was forcibly administered.

Accordingly, considering the results shown in Example 5
in combination, it was revealed that the specific components
of the indigenous microbiota were highly likely to belong to
the Gram-positive group, and that the spore-forming fraction
played an important role in the induction of Treg cells.

EXAMPLE 7

Next, the species of the intestinal microbiota which induced the accumulation of Treg cells in the colon as
suggested in Examples 4 to 6 were identified. Specifically, segmented filamentous bacteria (SFB), 16 strains of the
Bacteroides spp. (Bactero, 6 strains of B. vulgatus, 7 of the B. acidificiens group 1, and 3 of the B. acidificiens group
2), 3 strains of the Lactobacillus (Lacto. (L. acidophilus, L. fermentum, and L. murinum)), and 46 strains of Clostridium
spp. (Clost., refer to “Itok K., and Mitsuoka, T. Character-
ization of clostridia isolated from feces of limited flora
mouse and their effect on caecal size when associated with
germ-free mice. Lab. Animals 19: 111-118 (1985)”), or
microbiota collected from mice (SPF) bred under a conven-
tional environment was orally administered to GF-Balb/c
mice or GF-IQI mice. The mice were maintained in vinyl
isolators for three weeks. Then, CD4 cells were isolated from
the colon and the small intestine of these mice. The
numbers of Treg cells in the colon and the small intestine were
analyzed by flow cytometry.

The bacteria belonging to the genus Clostridium are
classified by sequencing of 16S rRNA gene, as follows.
Specifically, the 16S rRNA genes of the bacteria were
amplified by PCR using 16S rRNA gene-specific primer
pairs: 5'-AGAGTTTGATCMTGGCTCAG-3' (SEQ ID NO: 60)
and 5'-ATTACCGCGGTGCTG-3' (SEQ ID NO: 61)
(see T. Aebischer et al., Vaccination prevents Helicobacter
pylori-induced alterations of the gastric flora in mice. FEMS
PCR product was then introduced into pCR-Blnunt Vector.
The inserted were sequenced and aligned using the ChistalW
software program. The resulting sequences of 16S rRNA
genes derived from strain 1-41 of 46 strains of Clostridium
spp. were shown in SEQ ID NO: 21-61. A phylogenetic tree
was constructed by the neighbor-joining method with the
resulting sequences of the 41 strains of Clostridium and
those of known bacteria obtained from Genbank database
using Mega software.

The results showed no effect on the number of the Treg
cells in the colon was observed in the GF mice in which the
segmented filamentous bacteria (SFB) were colonized.
Moreover, mice in which the cocktail of three strains of
Lactobacillus was colonized gave similar results. On the
other hand, it was shown that the accumulation of Foxp3+ cells
in the colonic lamina propria was strongly induced in the
mice in which 46 strains of Clostridium spp. were
colonized. Importantly, such accumulation was promoted
irrespective of the genetic backgrounds of the mice, and led
to the increase in number similar to that in the SPF mice
although intestinal microbiota of only a single genus were
colonized. It was also shown that the colonization of the
Clostridium did not change the number of Treg cells in the
small intestinal lamina propria. Note that, when the 16
strains of Bactericides spp. were colonized, the number of
Treg cells in the colon was increased significantly. However,
the extent of the increase varied depending on the genetic
background of the mice in which the bacteria were
colonized.

EXAMPLE 8

Next, CD4 expression, Foxp3 expression, and Helios
expression in LP lymphocytes of the thymuses and the
colon of SPF mice, GF mice, Lactobacillus-colonized mice,
and Clostridium-colonized mice were analyzed by
flow cytometry.

The results show that most Foxp3+ cells found in the SPF
mice or the Clostridium-colonized mice did not express
Helios. Note that Helios is a transcription factor known to be
expressed in thymic-derived natural Treg cells (see A. M.
Thornton et al., J Immunol 184, 3433 (Apr. 1, 2010). Accordingly, it was suggested that most of the Treg cells in the SPF mice and the Clostridium-colonized mice were Treg cells induced in peripheral portions (so-called iTreg cells).

**EXAMPLE 9**

Next, it was investigated whether or not the colonization of the Clostridium or the like had an influence on other T cells. Specifically, SFB, 16 strains of Bacteroides spp. (Bacterio.), 46 strains of Clostridium spp. (Clost.), or microbiota collected from mice bred under a conventional environment (SPF) was colonized in GF Iqi mice. Three weeks later, lymphocytes in the colonic lamina propria were isolated from these mice, and stimulated with PMA (50 ng/ml) and ionomycin (1 μg/ml) for four hours in the presence of Golgistem (BD Bioscience). After the stimulation was given, intracellular cytokines were stained by using an anti-IL-17 PE antibody (TC11-18H10) and an anti-IFN-γ FITC antibody (BD Bioscience) in accordance with the manual of a cytokin/cytoplasm kit (BD Bioscience). Then, the ratio of IFN-γ+ cells or IL-17+ cells in CD4+ lymphocytes was analyzed by flow cytometry. The results show that the colonization of the Clostridium did not have any influence on TH1 cells (CD4+ IFN-γ+ cells) in the colon, and caused only a slight increase of TH17 cells (CD4+ IL-17+ cells). Accordingly, it was suggested that the genus Clostridium was a genus of bacteria which specifically induced Treg cells.

**EXAMPLE 10**

It has been reported that 46 strains of Clostridium spp. exert an influence on the accumulation of CD8+ intestinal tract intraepithelial lymphocytes (IELs) in the colon. Accordingly, it is conceivable that Clostridium regulates the immune system in various aspects, and that Clostridium exhibits a marked ability to induce and maintain Treg cells especially in the colon, as described above. In addition, a kind of cytokines, transforming growth factor-β (TGF-β), is known to play an important role in regulation of Treg cell generation.

In this respect, it was examined whether or not the colonization of Clostridium provided a colonic environment rich in TGF-β. Specifically, first, the whole colonos of GF mice, Clostridium-colonized mice, and Lactobacillus-colonized mice were cultured for 24 hours, and the culture supernatants thereof were measured for the concentration of active TGF-β (TGF-β1) by ELISA (the number of mice analyzed was four per group).

The results show that the amount of TGF-β produced in the colon of the Clostridium-colonized mice was significantly greater than that in the colon of GF mice and the Lactobacillus-colonized mice.

Next, intestinal epithelial cells (IECs) of GF mice and Clostridium-colonized mice were cultured for 24 hours, and the culture supernatants thereof were measured for the concentration of active TGF-β (TGF-β1) by ELISA (the number of mice analyzed was four per group).

The results show that TGF-β was detected in the culture supernatant of the IECs isolated from the Clostridium-colonized mice, whereas no TGF-β was detected in the culture supernatant of the IECs isolated from the GF mice.

Next, as described above, splenic CD4+ T cells were cultured for five days together with a 50% conditioned medium in which IECs isolated from the GF mice or the Clostridium-colonized mice were cultured, and with the anti-CD3 antibody, in the presence or absence of an anti-TGF-β antibody. Then, the T cells were collected, and analyzed for expression of Foxp3 by real-time RT-PCR.

The results show that when the culture supernatant of the IECs derived from the Clostridium-colonized mice was added to the splenic CD4+ T cells, differentiation into Foxp3-expressing cells was accelerated. Meanwhile, differentiation into Treg cells was inhibited by the anti-TGF-β antibody.

The expression of MMP2, MMP9, and MMP13, which are thought to contribute to the activation of latent TGF-β was investigated. The expression of indoleamine 2,3-dioxygenase (IDO), which is thought to be involved in the induction of Treg cells, was also investigated. Specifically, 46 bacterial strains of the genus Clostridium (Clost.), or three bacterial strains of the genus Lactobacillus (Lacto.) were orally administered to C57BL/6 germ-free mice. Three weeks after administration, IECs were collected, and analyzed for relative mRNA expression levels of MMP2, MMP9, MMP13, and IDO genes by real-time RT-PCR (the number of mice analyzed was three per group).

For the relationship between the activation of latent TGF-β and the above-described MMP, see D’Angelo et al., J. Biol. Chem. 276, 11347-11353, 2001; Heidinger et al., J. Biol. Chem. 387, 69-78, 2006; Yu et al., Genes Dev. 14, 163-176, 2000. For the relationship between IDO and the induction of Treg cells, see G. Mattei et al., Gut 59, 595 (May, 2010).

The results show in agreement with the production of TGF-β described above, that transcription products of the genes encoding MMP2, MMP9, and MMP13 were expressed at higher levels in the IECs derived from the Clostridium-colonized mice than in those in the GF mice and in the Lactobacillus-colonized mice.

Moreover, IDO was expressed only in the Clostridium-colonized mice.

Accordingly, it was revealed that the Clostridium activated the IECs, and led to the production of TGF-β and other Treg cell-inducing molecules in the colon.

**EXAMPLE 11**

Next, it was investigated whether or not the Treg cell accumulation induced by the colonization of the Clostridium was dependent on signal transmission by pathogen-associated molecular pattern recognition receptors. Specifically, the numbers of Treg cells in the colonic lamina propria of each SPF mice of Myd88−/− (deficient in Myd88 signaling adaptor for Toll-like receptor), Rip2−/− (deficient in Rip2 (NOD receptor adaptor)), and Il1R1−/− (deficient, essential signal transmission factor for Dectin-1 signal transmission) were examined. In addition, Clostridium spp. were caused to be colonized in the Myd88−/− GF mice, and the change in the number of Treg cells was investigated. The results show that the number of Treg cells of each kind of the SPF mice deficient in the associated factors of the pathogen-associated molecular pattern recognition receptors did not change relative to that of wild-type mice of the same litter, which served as a control. In addition, it was found that when Clostridium spp. were colonized in GF mice deficient in Myd88, the accumulation of Treg cells in the colonic lamina propria was induced. Accordingly, it has been suggested that the mechanism of inducing the accumulation of Treg cells in the colonic lamina propria relies on activation of recognition pathway for major pathogen-associated molecular patterns as is caused by most bacteria, but on specific commensal bacterial species.

**EXAMPLE 12**

Intestinal tract Foxp3+ Treg cells are known to exert some immunosuppressive functions through IL-10 production (re-
fer to NPL 9). Meanwhile, animals having CD4⁺ Foxp3⁺ cells from which IL-10 is specifically removed are known to develop inflammatory bowel disease (refer to NPL 18). In this respect, first, the expression of IL-10 in lymphocytes of various tissues was examined. Specifically, lymphocytes were isolated from various tissues of SPF II10⁻/⁻ mice, and the expression of CD4 and the expression of Venus were analyzed by flow cytometry.

Lymphocytes in the colonic lamina propria were isolated from II10⁻/⁻ mice, and the expression of T cell receptor β chain (TCRβ) on the surfaces of the cells was detected by FACS.

Lymphocytes in the colonic lamina propria were isolated from II10⁻/⁻ mice. The lymphocytes were stimulated with PMA (50 ng/ml) and ionomycin (1 μg/ml) for four hours in the presence of Golgistop (BD Bioscience). Then, after the stimulation was given, intracellular cytokines were stained by using an anti-IL-17 PE antibody, an anti-IL-4 APC antibody (11B11), and an anti-IFN-γ FITC antibody (BD Bioscience) in accordance with the manual of a cytokine/cytokine kit (BD Bioscience).

In addition, Foxp3⁺ CD4⁺ cells and Foxp3⁻ CD4⁺ cells were isolated from the spleen (Spl) of Foxp3⁺GFP reporter mice, and Venus⁺ cells were isolated from the colonic lamina propria and the small intestine (SI) lamina propria of II10⁻/⁻ mice. The obtained cells were analyzed in terms of expression of predetermined genes. The gene expression was analyzed by real-time RT-PCR using a Power SYBR Green PCR Master Mix (Applied Biosystems) and an ABI 7300 real-time PCR system (Applied Biosystems). Here, the value for each cell was normalized for the amount of GAPDH.

The results show that almost no Venus⁺ cells (IL-10-producing cells) were detected in the cervical lymph nodes (peripheral lymph nodes), thymus, peripheral blood, lung, and liver of mice kept under the SPF conditions. Meanwhile, in the spleen, Peyer’s patches, and mesenteric lymph nodes thereof, Venus⁺ cells were slightly detected. On the other hand, many Venus⁺ cells were found in the lymphocytes in the small intestine lamina propria and colonic lamina propria. In addition, most of the Venus⁺ cells in the intestines were positive for CD4, and also positive for T cell receptor β chain (TCRβ). It was found that the Venus⁺ CD4⁺ T cells and CD4⁺ T cells expressed Foxp3 and other Treg cell-associated factors such as a cytotoxic T-Lymphocyte antigen (CTLA-4) and a glucocorticoid-induced TNFR-associated protein (GITR), although the Venus⁺ CD4⁺ T cells showed none of the phenotypes of Th2 (IL-4-producing) and Th17 (IL-17-producing). It was shown that the expression level of CTLA-4 in the intestinal Venus⁺ cells was higher than that in the splenic GFP⁺ Treg cells isolated from the Foxp3⁺GFP reporter mice.

**EXAMPLE 13**

Venus⁺ cells can be classified into at least two subsets, namely, Venus⁺ Foxp3⁺ double positive (DP) Treg cells and Venus⁺ Foxp3⁻ Treg cells on the basis of intracellular Foxp3 expression. Cells of the latter subset correspond to type 1 regulatory T cells (Tr1) (refer to NPL 8 and 9). In this respect, the Venus⁺ cells (IL-10-producing cells) observed in Example 8 were investigated in terms of the expression of Foxp3. Specifically, the expression of CD4, Foxp3, and Venus in the lamina propria of the colon and the lamina propria of the small intestine of II10⁻/⁻ mice kept under GF or SPF conditions was analyzed by FACS, and the numbers of Venus⁺ cells in the intestinal tract lamina propria were compared between SPF and GF II10⁻/⁻ mice.

In addition, the intracellular expression of Venus and Foxp3 in CD4 cells in various tissues of SPF II10⁻/⁻ mice was analyzed by flow cytometry.

In order to investigate whether or not the presence of commensal bacteria had any influence on the expression of IL-10 in regulatory cells in the gastrointestinal tracts, germ-free (GF) II10⁻/⁻ mice were prepared. Then, predetermined species of bacteria were caused to be colonized in the obtained GF II10⁻/⁻ mice. Three weeks after the species of bacteria were colonized, a CD4⁺ cell group (V⁺F⁺, Venus⁺ Foxp3⁺ cells; V⁺F⁻, Venus⁻ Foxp3⁻ cells; and V⁻F⁺, Venus⁻ Foxp3⁺ cells) in which Foxp3 and/or Venus were expressed in the colon and the small intestine was analyzed by flow cytometry.

In order to check whether or not the presence of commensal bacteria had any influence on the expression of IL-10 in regulatory cells in the gastrointestinal tracts, antibiotics were orally given with water to five or six II10⁻/⁻ mice per group for 10 weeks. The following antibiotics were used in combination: ampicillin (A; 500 mg/L Sigma), vancomycin (V; 500 mg/L NACALAI TESQUE, INC.), metronidazole (M; 1 g/L NACALAI TESQUE, INC.), and neomycin (N; 1 g/L NACALAI TESQUE, INC.).

Then, CD4 and Foxp3 of lymphocytes in the lamina propria of the colon, the lamina propria of the small intestine (SI), and Peyer’s patches were stained with antibodies, and analyzed by FACS.

The results were obtained from two or more independent experiments which gave similar results.

The results show that the small intestinal lamina propria was rich in Venus⁺ Foxp3⁺ cells, namely, Tr1-like cells, and that the Venus⁺ Foxp3⁺ DP Treg cells were present at a high frequency in the colon of the SPF mice. In contrast, although sufficient numbers of Foxp3⁺ cells were observed also in other tissues, the expression of Venus⁺ cells was not observed in almost all of the cells.

In addition, it was shown that all regulatory T cell fractions of Venus⁺ Foxp3⁺, Venus⁺ Foxp3⁻, and Venus⁻ Foxp3⁺ in the colon significantly decreased under the GF conditions. Moreover, similar decrease in Venus⁺ cells was observed also in the SPF II10⁻/⁻ mice treated with the antibiotics.

The colonization of *Clostridium* spp. strongly induced all regulatory T cell fractions of Venus⁺ Foxp3⁺, Venus⁺ Foxp3⁻, and Venus⁻ Foxp3⁺ in the colon, and the degrees of the induction thereof were equal to those in the SPF mice. In addition, it was found that the colonization of the three strains of *Lactobacillus* or the colonization of SFB had an extremely small influence on the number of Venus⁺ and/or Foxp3⁺ cells in the colon. Moreover, the colonization of 16 strains of *Bacteroides* spp. also induced Venus⁺ cells, but the influence of the colonization was specific to Venus⁺ Foxp3⁺ Tr1-like cells. On the other hand, it was found that none of the bacterial species tested exerted any significant influence on the number of IL-10-producing cells in the small intestinal lamina propria (refer to FIG. 26).

Hence, it was shown that the genus *Clostridium* colonized in the colon or a physiologically active substance derived from the bacteria provided a signal for inducing the accumulation of IL-10 regulatory T cells in the colonic lamina propria or the expression of IL-10 in T cells. It was shown that the number of Venus⁺ cells in the small intestine was not significantly influenced by the situation where no commensal bacteria were present or commensal bacteria were
decreased, and that IL-10+ regulatory cells (Tr1-like cells) accumulated in the small intestinal lamina propria independently of commensal bacteria.

EXAMPLE 14

It was investigated whether or not Venus+ cells induced by the genus Clostridium had an immunosuppressive function similar to that of Venus+ cells in the colon of SPF mice. Specifically, CD4+ CD25+ cells (effector T cells, Teff cells) isolated from the spleen were seeded in a flat-bottomed 96-well plate at 2 x 10^4/well, and cultured for three days together with 2 x 10^4 splenic CD11c+ cells (anti-gen-representing cells) subjected to 30 Gy radiation irradiation treatment, 0.5 μg/ml of an anti-CD3 antibody, and a lot of Treg cells. In addition, for the last six hours, the CD4+ CD25+ cells were cultured with [3H]-thymidine (1 μCi/well) was added thereto. Note that, Teff cells used in Example 14 were CD4+ GFP+ T cells isolated from the spleen of Foxp3+GFp reporter mice, or CD4+ Venus+ T cells in the colonic lamina propria of GF110venus mice in which Clostridium spp. were colonized or SPF110venus mice. Then, proliferation of the cells was determined based on the uptake amount of [3H]-thymidine, and represented by a count per minute (cpm) value.

The results show that Venus+ CD4+ cells of the mice in which the genus Clostridium was colonized suppressed in vitro proliferation of CD25+ CD4+ activated T cells. The suppression activity was slightly inferior to that of GFP+ cells isolated from the Foxp3+GFp reporter mice, but equal to that of Venus+ cells isolated from the SPF110venus mice. Accordingly, it has been shown that the genus Clostridium induces IL-10-expressing T cells having sufficient immunosuppressive activities, and thereby plays a critical role in maintaining immune homeostasis in the colon.

EXAMPLE 15

Next, the influence of the colonization of a large number of Clostridium on the local immune response and the resultant proliferation of Treg cells was investigated.

Dextran Sulfate Sodium (DSS)-Induced Colitis Model>

First, the DSS-induced colitis model was prepared as described above, and the influence on the model mice of the inoculation of Clostridium and the proliferation of Treg cells was investigated. Specifically, control mice and Clostridium-inoculated mice were treated with 2% DSS, then observed and measured for six days for body weight loss, the hardness of stool, and bleeding, and then were evaluated numerically. In addition, on day 6, the colons were collected, dissected, and analyzed histologically by H&E staining.

The results show that the symptoms of the colitis such as body weight loss and rectal bleeding were significantly suppressed in the mice having a large number of Clostridium (hereinafter also referred to as “Clostridium-abundant mice”) in comparison with the control mice (C57BL/6 mice grown in a conventional environment for six weeks and not inoculated with the fecal suspension). All the features typical for colonic inflammation, such as shortening of the colon, edema, and hemorrhage, were observed markedly in the control mice in comparison with the Clostridium-abundant mice. Moreover, histological features such as mucosal erosion, edema, cellular infiltration, and crypt loss were less severe in the DSS-treated Clostridium-abundant mice than in the control mice.

OXAZOLONE-INDUCED COLITIS MODEL

Next, the oxazolone-induced colitis model was prepared as described above, and the influence on the model mice of the inoculation of Clostridium and the proliferation of Treg cells was investigated. Specifically, control mice and Clostridium-inoculated mice were sensitized with oxazolone, and subsequently the inside of the rectums thereof were treated with a 1% oxazolone/50% ethanol solution. Then, body weight loss was observed and measured. In addition, the colons were dissected, and analyzed histologically by H&E staining.

The results show that the colitis proceeded along with persistent body weight loss in the control mice. Meanwhile, the body weight loss of the Clostridium-abundant mice was reduced. In addition, it was also revealed that portions having histological diseases such as mucosal erosion, edema, cellular infiltration, and hemorrhage were reduced in the colon of the Clostridium-abundant mice.

EXAMPLE 16

Next, the influence, on the systemic immune response (systemic IgE production), of the colonization of a large number of Clostridium and the resultant proliferation of T cells was investigated. Specifically, as described above, control mice and Clostridium-inoculated mice were immunized by administering alum-absorbed ovalbumin (OVA) twice at a 2-week interval. Then, sera were collected from these mice, and the OVA-specific IgE level thereof was investigated by ELISA. In addition, splenic cells were collected from the mice in each group, and IL-4 and IL-10 production by in-vitro OVA restimulation was investigated.

Results show that the IgE level was significantly lower in the Clostridium-abundant mice than in the control mice. Moreover, the IL-4 production by the OVA restimulation was reduced and the IL-10 production thereby was increased in the splenic cells of the Clostridium-abundant mice sensitized with OVA and alum, in comparison with those of the control mice.

Accordingly, in consideration of the results shown in Example 15 in combination, the induction of Treg cells by Clostridium in the colon plays an important role in both local and systemic immune responses.

EXAMPLE 17

Next, GF Balb/c were colonized with three strains of Clostridium belonging to cluster IV (strains 22, 23 and 32 listed in FIG. 49). Three weeks later, colonic Foxp3+ Treg cells were analyzed by FACS. Results show that gnotobiotic mice colonized with three strains of Clostridium showed an intermediate pattern of Treg induction between GF mice and mice inoculated with all 46 strains.

EXAMPLE 18

Next, it was investigated whether or not a spore-forming (for example, a chloroform resistant) fraction of a fecal sample obtained from humans had the effect of inducing proliferation or accumulation of regulatory T cells similar to the spore-forming fraction of the fecal sample obtained from mice.

Human stool from a healthy volunteer (Japanese, male, 29 years old) was suspended with phosphate-buffered saline (PBS), mixed with chloroform (final concentration 3%), and then incubated in a shaking water bath for 60 min. After evaporation of chloroform by bubbling with N2 gas, the aliquots containing chloroform-resistant (for example,
spore-forming) fraction of human intestinal bacteria were orally inoculated into germ-free (GF) mice (IQI, 8 weeks old). The treated mice were kept in a vinyl isolator for 3 weeks. The colon was collected and opened longitudinally, washed to remove fecal content, and shaken in Hanks' balanced salt solution (HBSS) containing 5 mM EDTA for 20 min at 37°C. After removing epithelial cells and fat tissue, the colon was cut into small pieces and incubated with RPMI-1640 containing 4% fetal bovine serum, 1 mg/ml collagenase D, 0.5 mg/ml dispase and 40 μg/ml DNase I (all manufactured by Roche Diagnostics) for 1 hour at 37°C in a shaking water bath. The digested tissue was washed with HBSS containing 5 mM EDTA, resuspended in 5 ml of 40% Percoll (manufactured by GE Healthcare) and overlaid on 2.5 ml of 80% Percoll in a 15-ml Falcon tube. Percoll gradient separation was performed by centrifugation at 780 g for 20 min at 25°C. The interface cells were collected and suspended in staining buffer containing PBS, 2% FBS, 2 mM EDTA and 0.09% NaN3 and stained for surface CD4 with Phycoerythrin-labeled anti-CD4 Ab (RM4-5, manufactured by BD Biosciences). Intracellular staining of Foxp3 was performed using the Alexa647-labeled anti-FoxP3 Ab (E13-1656, manufactured by eBioscience) and Foxp3 Staining Buffer Set (manufactured by eBioscience). The percentage of Foxp3 positive cells within the CD4 positive lymphocyte population was analyzed by flow cytometry.

Results show that when the spore-forming (for example, the chlorosome resistant) fraction of human intestinal bacteria was colonized in GF mice, the accumulation of Foxp3+ regulatory (Treg) cells in the colonic lamina propria of the mice was induced.

Next, it was investigated what species of bacteria grew by gavaging with chlorofom-treated human stool.

Specifically, using a QIAamp DNA Stool mini kit (manufactured by QIAGEN), bacterial genomic DNA was isolated from the healthy volunteer as described above (human stool) or fecal pellets from GF mice gavaged with chlorofom-treated human stool (GF+Chloro.). Quantitative PCR analysis was carried out using a LightCycler 480 (manufactured by Roche). Relative quantity was calculated by the ΔCt method and normalized to the amount of total bacteria, dilution, and weight of the sample. The following primer sets were used:

<table>
<thead>
<tr>
<th>Primer Set</th>
<th>SEQ ID NO.</th>
</tr>
</thead>
<tbody>
<tr>
<td>total bacteria</td>
<td>62</td>
</tr>
<tr>
<td>5'-GGTGAATAGTCTGCCG-3'</td>
<td>63</td>
</tr>
<tr>
<td>5'-TACCCATCTCTCTCTG-3'</td>
<td>64</td>
</tr>
<tr>
<td>Clostridium cluster XIVA (Clostridium cocoides subgroup)</td>
<td>65</td>
</tr>
<tr>
<td>5'-AATGACCCGACCTGACTA-3'</td>
<td>66</td>
</tr>
<tr>
<td>5'-CTTACGGTTTCTCTTGCAA-3'</td>
<td>67</td>
</tr>
<tr>
<td>Clostridium cluster IV (Clostridium leptum)</td>
<td>68</td>
</tr>
<tr>
<td>5'-GGCATGAGCTGAGT-3'</td>
<td>69</td>
</tr>
<tr>
<td>5'-CTTTCTCCCTTGTTTCGAA-3'</td>
<td>70</td>
</tr>
</tbody>
</table>

Results show that gavaged with chlorofom-treated human stool had large amounts of spore-forming bacteria, such as Clostridium clusters XIVA and IV, and a severe decrease of non-spore-forming bacteria, such as Bacteroides, compared with the human stool before chlorofom treatment.

**EXAMPLE 19**

Human stool (2 g) from a healthy volunteer (Japanese, male, 29 y old) was suspended with 20 ml phosphate-buffered saline (PBS), mixed with or without chlorofom (final concentration 3%), and incubated in a shaking water bath for 60 min. The chlorofom was then evaporated by bubbling with N2 gas for 30 min. The suspensions of untreated human feces (designated as 'huUT') and chlorofom-treated human feces (designated as 'huChloro') were orally inoculated into Germ-free (GF) mice (IQI, 8 week old) (250 µl/mouse). The suspension of huUT was inoculated into 4 GF mice, which were numbered from #A1 to #A4, and that of huChloro was inoculated into 4 GF mice numbered from #B1 to #B4. Such GF mice which were inoculated with suspensions of feces or the like are also referred to as “ex-GF mice” hereinafter. Each group of ex-GF mice was separately kept in a vinyl isolator to avoid further microbial contamination. After 3 weeks, the small intestinal and colonic lamina propria lymphocytes from each mouse were separately collected, and examined for the expressions of surface CD4 and intracellular Foxp3, Helios, IL-17 and IFN-γ by flow cytometry. For intracellular IL-17 and IFN-γ staining, isolated lymphocytes were stimulated in vitro with PMA and ionomycin for 4 hours. Foxp3 is the transcription factor essential for the differentiation and function of Treg cells. Helios is a member of the Ikaros transcription factor family and Helios−Foxp3+ Treg cells have been suggested to be Treg cells induced in the periphery [so called induced Treg (iTreg) cells], as shown in FIGS. 1A-D, the percentages of Foxp3+ Treg cells within CD4+ T cells in the small intestinal and colonic lamina propria of both groups of ex-GF mice were increased, compared with those in GF mice. Marked increases were also observed for the percentage of Helios− cells among Foxp3+ Treg cells in small intestine and colon in both groups of ex-GF mice. Notably, besides Foxp3+ Treg cells, a significant accumulation of IL-17-expressing CD4+ cells (namely, Th17 cells) was observed in exGF+huUT mice, whereas it was only marginally observed in exGF+huChloro mice (FIGS. 1E, F). In both groups of mice, the percentages of IFN-γ+ cells in CD4+ cells were unchanged (FIGS. 1E, G).

**EXAMPLE 20**

To investigate whether dead bacteria also have an effect on the induction of Treg cells, the suspension of chlorofom-treated human feces was autoclaved (121°C for 20 min) and orally inoculated into GF mice (once a week for 4 weeks). After 4 weeks, mice were sacrificed, and the colonic lamina propria lymphocytes from each mouse were examined for the expression of CD4, Foxp3 and Helios by flow
cytometry. As shown in FIG. 2, the inoculation of dead bacteria exhibited no effect on the numbers of Foxp3+ cells or Helios-Foxp3+ cells. These results do not rule out the possibility that the amount of dead bacteria inoculated was not sufficient, but suggest that live bacteria are required for the induction of Treg cells.

EXAMPLE 21

To confirm the induction of Treg cells by chloroform-resistant bacteria, another stool was obtained from the same person on a different day, treated with chloroform, and inoculated into IQR GF mice (7 mice, numbered from #C1 to #C7). After 3-4 weeks, mice from #C1 to #C5 were sacrificed, and the small intestinal and colonic lamina propria lymphocytes from each mouse were separately collected, and examined for the expression of CD4 and Foxp3 by flow cytometry. Consistent with the findings in Example 19, colonization with chloroform-treated human feces significantly increased the accumulation of Foxp3+CD4+ Treg cells in colonic and small intestinal lamina propria (FIG. 3). These results further support the notion that chloroform-resistant spore-forming bacteria can induce differentiation, proliferation and/or recruitment of Treg cells in intestinal lamina propria.

EXAMPLE 22

To test whether Treg cell induction by chloroform-resistant spore-forming fraction of human intestinal bacteria is horizontally transmissible, IQR GF mice (6 mice, numbered from #D1 to #D6) were cohoused for 4 weeks with mice #C6 and #C7 in the same cage in a vinyl isolator. Lamina propria lymphocytes from colon and small intestine were isolated and examined for CD4 and Foxp3. Cohoused mice exhibited a significant increase in the percentage of Foxp3+ cells among CD4+ cells (FIG. 4). Therefore, Treg cell induction by human intestinal bacteria is horizontally transmissible. These results let us assume a role of prominent components of the intestinal microbiota, rather than minor components, for the induction of Treg cells.

EXAMPLE 23

The frozen stock of cecal content from mouse #C4 was thawed, suspended in 10 times its volume (w/v) of PBS, and passed through a 70 μm cell strainer. The suspension was then treated with 3% chloroform, diluted 2000- or 20000-fold with PBS, and orally inoculated into GF IQR mice (2.5×10^7 or 2.5×10^6 bacterial cells/250 μl/head, respectively). The 2000-fold diluted sample was orally inoculated into 4 mice (designated as exGF+2000, numbered from #E1 to #E4), whereas 20000-fold diluted samples was inoculated into 8 mice (designated as exGF+20000, numbered from #F1 to #F8). After 3 weeks, the intestinal lamina propria lymphocytes were isolated and examined for CD4, Foxp3 and Helios. Both 2000- and 20000-fold diluted samples similarly induced a marked accumulation of Foxp3+CD4+ cells in the intestinal lamina propria (FIG. 5). Therefore, the dose of bacteria for oral inoculation can be minimized to less than 2.5×10^6 bacterial cells.

EXAMPLE 24

The frozen stock of cecal content from mouse #F3, #F7 or #F8 was suspended in 10 times its volume (w/v) of PBS, passed through a 70 μm cell strainer, and treated with 3% chloroform. Then, the fecal suspension from mouse #F3 was orally inoculated into 5 GF mice (numbered from #G1 to #G5), that from #F7 mouse into 4 GF mice (numbered from #H1 to #H4), and that from #F8 mouse into 4 GF mice (numbered from #I1 to #I4). After 4 weeks, lymphocytes from colonic and small intestinal lamina propria were isolated and examined for CD4, Foxp3 and Helios expression by flow cytometry. All #F#, #G# and #H# mice exhibited a significant increase in the percentage of Foxp3+ cells among CD4+ cells in the intestinal lamina propria compared with untreated GF mice (FIG. 6). Therefore, the Treg cell induction by human intestinal bacteria colonizing in exGF+20000 mice is also transmissible. Moreover, as shown in the later meta 16S rDNA sequencing data (FIG. 8), these mice commonly had bacteria having 16S rDNA sequence similarities with 16S rDNA sequence similarities with 20 species of known bacteria (C. oryzae, H. succinivorans, E. fischeri, H. filiformis, C. clostridioforme, C. indolis, C. bolteae, Bacteroides sp. MANG, L. bacterium DJ-VP90, Ruminococcus sp. ID8, C. lavalense, C. symbiosum, E. consortum, C. saccharogena, C. ramosum, Flavonifractor plautii, C. scindens, Clostridium sp. 2335, Clostridium sp. 14616 and Clostridium sp. MLG055).

EXAMPLE 25

A frozen stock of the cecal content from #F8 mouse was serially diluted with 0.85% NaCl under an aerobic condition and plated onto BL agar. After growth at 37°C for 2 or 4 days, 50 colonies were observed. Of the 50 colonies, 29 were picked up, cultured for additional 2 or 4 days at 37°C, and stored in EG stock medium (10% DMSO) at −80°C. The genomic DNA from each colony was isolated, and 16S rRNA coding gene sequence was analyzed. The sequence of 16S rRNA of each colony revealed that the 29 colonies observed were represented by three strains, each having 100% similarity with Clostridium ramosum, 93.75% with Clostridium saccharogena, 100% with Flavonifractor plautii, 99.17% with Clostridium hathewayi, 99.23% with Clostridium scindens, or 99.68% with Clostridium sp. 2335. Within the 29 colonies that were selected from the original 50 colonies, only Clostridium saccharogena, Clostridium ramosum, and Flavonifractor plautii were present (25, 3, and 1 colonies, respectively). These 3 isolated strains were propagated, mixed and inoculated into GF IQR mice (4 mice, numbered from #J1 to #J4). After 3-4 weeks, the colonic lamina propria lymphocytes were collected, and examined for the expression of CD4, Foxp3, and Helios by flow cytometry. Foxp3+ cells or Helios+ cells were not induced or only weakly induced by the colonization of these strains of bacteria in the colon (FIG. 7). These results suggest that the combination of Clostridium saccharogena and Clostridium ramosum (both within cluster XVIII) were insufficient to induce Treg cells in the colon of mice. The effects of Flavonifractor plautii were not clear, since the strain was only represented by 1 of the 29 colonies that were selected.

EXAMPLE 26

The frozen glycerol stock of cecal content from #G2 mouse was suspended with PBS, seeded onto BL agar plate, and incubated for 48 hours, similarly to the procedure done in Example 19. Different from Example 19, all bacteria on the plate were collected by scraping with a plate scraper, suspended in TS broth and inoculated into GF IQR mice (4 mice, numbering from #K1 to #K4). It should be noted that
the bacterial suspension used in this experiment included bacteria that did not propagate but survived on the plate. After 4 weeks, lamina propria lymphocytes from colon and small intestine of K1–K4 mice were isolated and examined for CD4, Foxp3 and Helios expression. All 4 mice exhibited a significant increase in the percentages of Foxp3+ cells among CD4+ cells (FIGS. 9A, 9B) and Helios+ cells among Foxp3+ Treg cells (FIGS. 9A, 9C) in the intestinal lamina propria compared with untreated GF mice. Considering that the inoculation of mice with 6 strains of bacteria propagated on the BL agar plate failed to induce Treg cells, bacteria that did not propagate but survived on the plate might be responsible for the induction of Treg cells.

EXAMPLE 27

Bacterial DNA was extracted from the cecal contents of mouse #A1, #C4, #F2, #G2, #I3, #J3 and #K3. Variable region 1-2 (V1-2) in bacterial 16S rRNA coding gene were amplified by PCR and used as template for metasequencing. Resulting sequences (3400 reads for each sample) were classified into operational taxonomic units (OTUs) on the basis of sequence similarity (>97% identity). Representative sequences from each OTU were compared with sequences in nucleic acid databases using BLAST to determine their closest relatives in known species. The numbers of detected reads and the closest relatives for each OTU are shown in Table 1. The relative abundances of OTUs having the same closest relative in each cecal sample are shown in FIG. 8. In mouse #A1, 153 OTUs (their closest relatives were 93 species) were identified and half of them were related to Bacteroides species. In contrast, in mouse #C4, 113 OTUs were identified and most of them were related to species belonging to the family Clostridiaceae. In mouse #F2, #G2, #I3, #J3 and #K3, 97-68 OTUs were identified. In these mice, in which Treg cell accumulation was observed in the intestine, the majority of bacteria consisted of bacteria having 16S rDNA sequence similarities with C. amniliicum, H. saccharovorans, E. fissaicata, H. filiformis, C. clostridiforme, C. indolis, C. bolteae, Bacteroides sp. MANG, L. bacterium DFJ_VP30, Ruminococcus sp. ID8, C. lavalense, C. symbiosum, E. contortum, C. saccharogranum, C. ramosum, P. platii, C. scindens, Clostridium sp. 2335, Clostridium sp. 14616 and cf Clostridium sp. MLG055.

In mouse #J3, in which Treg cell accumulation was not observed, 3 OTUs were detected. Each has the 16S rDNA sequence similarity with C. saccharogranum, C. ramosum or F. platii. These results suggest that the combination of these three species may be insufficient to induce the intestinal Treg cells accumulation.

EXAMPLE 28

Bacterial strains were isolated from the cecal contents of mouse #F2, #G2, #I1 and #K using BL agar or EG agar plates. Applicant picked-up 144 colonies from EG agar plates and 116 colonies from BL agar plates. BLAST search of 16S rRNA coding sequence of these clones revealed that they belonged to 17 species, and each had 93-100% similarities with C. indolis, C. bolteae, Bacteroides sp. MANG, L. bacterium DFJ_VP30, A. coliforminis, Ruminococcus sp. ID8, C. lavalense, C. symbiosum, E. contortum, C. saccharogranum, C. ramosum, P. platii, C. hathewayi, C. scindens, Clostridium sp. 2335, Clostridium sp. 14616 and cf Clostridium sp. MLG055 (Table 2). They all belonged to Clostridium clusters IV, XIVA or XVIII (2 species of cluster IV, 12 of cluster XIVA, 1 of cluster XVI and 2 of cluster XVIII).

EXAMPLE 29

Of the colonies selected in Example 28, additional colonies were picked and isolated and these strains were cultured using EG and BL media. BLAST search of 16S rRNA coding sequence of these clones revealed that they belonged to a total of 31 species (including the species mentioned in Example 28), and each had 93-100% similarities with Clostridium saccharogranum, Clostridium ramosum, Pseudoflavonifractor capillosus ATCC 29799, Clostridium hathewayi, Clostridium saccharolyticum WM1, Bacteroides sp. MANG, Clostridium saccharolyticum, Clostridium scindens, Lachnospiraceae bacterium 5_1_57F AA, Lachnospiraceae bacterium 6_1_63 F AA, Clostridium sp. 14616, Clostridium bolteae ATCC BAA-613, cf. Clostridium sp. MLG055, Erysipelotrichaceae bacterium 2_2_44A, Clostridium indolis, Anaerostipes caceae, Clostridium bolteae, Lachnospiraceae bacterium DJF VP30, Lachnospiraceae bacterium 3_1_57F AA CT1, Anaerotrunctus colihomnis, Anaerotruncus colihomnis DSM 17241, Ruminococcus sp. ID3, Lachnospiraceae bacterium 2_1_46F AA, Clostridium lavalense, Clostridium asporagiforme DSM 15981, Clostridium symbiosum, Clostridium symbiosum WAI-14163, Eidberarterium contortum, Clostridium sp. D5, Oscillospiraceae bacterium NML 061048, Oscillibacter valericigenes, Lachnospiraceae bacterium A4, Clostridium sp. 316002/08, and Clostridiales bacterium 1_7_47F AA, Blautia cocoides, Anaerostipes caceae DSM 14662 (Table 3). The stocks of bacterial strains were stored in 10% glycerol stock plus the media used to grow the cultures, and tubes were stored in a –80°C freezer.

EXAMPLE 30

To investigate whether the strains in Example 29 have the ability to induce Tregs in GF mice, 31 strains on Table 3 were mixed at equal amounts of media volume using TS media and inoculated into GF mice. A detailed analysis of the 16S rRNA sequences revealed that 8 of the 31 strains overlapped with other strains (see Table 3, indicated by an asterisk), resulting in 23 distinct bacterial strains. As shown in FIG. 10, when orally administered to GF mice, the mixture of the 23 strains (23 mix) induced very strong levels of Tregs (35-40% in the colon lamina propria, >10% in the small intestine; FIG. 10). These Tregs observed with colonic by 23 mix were mostly Helios+.

EXAMPLE 31

To investigate whether the abundant members of the intestinal microbiota in the chloriform-resistant fraction of human intestinal bacteria, rather than the minor members, drive the induction of Treg cells, adult GF mice were inoculated with diluted caecal samples from mice that had been inoculated with the chloriform-resistant fraction of human intestinal bacteria (+hucChlo mice) as described in example 19. As shown in FIG. 11, even when the hucChlo mice caecal samples were diluted (diluted 2×10^5 and 2×10^6) to create 2×10^5 mice and 2×10^5 mice respectively, Tregs were induced in these adult GF mice.

EXAMPLE 32

To investigate whether the mix of 23 strains in Example 30 has the ability to induce Tregs in adult GF IQI mice more
effectively than *Faecalibacterium prausnitzii*, a well-known human Clostridia strain characterized for enhancing regulatory cell functions. 23 strains in table 4 were mixed in equal amounts with media to make a cocktail, which was then administered to adult IQuI GF mice. For comparison, *Faecalibacterium prausnitzii* was administered to another group of IQuI GF mice. As shown in FIG. 12, when orally administered to adult IQuI GF mice, the mixture of the 23 strains (23-mix) induced higher levels of Tregs than *Faecalibacterium prausnitzii*. *Faecalibacterium prausnitzii* (+Faecali.) showed negligible levels of Treg induction.

**EXAMPLE 33**

To investigate whether the microbiota communities in the +2x10^4 mice, described in example 31, were stable, serial oral inoculation of adult GF mice was performed to create +2x10^4-re mice (secondary inoculation) and +2x10^4-re-re (tertiary inoculation). As shown in FIG. 13 there was significant induction of Tregs in both the +2x10^4-re mice and the +2x10^4-re-re mice. To further eliminate nonessential components of the microbiota for Treg cell induction, the caecal content of +2x10^4 mice, described in example 31, was again diluted 2x10^4-fold and orally inoculated into another set of adult GF mice (+2x10^4^2 mice). As shown in FIG. 13, the (+2x10^4^2 mice exhibited a marked accumulation of Treg cells in the colon.

**EXAMPLE 34**

To assess the composition of the gut microbiota in +huUT (+hu), +huChlo, +2x10^8, +2x10^8-re and (+2x10^8), described in example 19, example 31, and example 33, bacterial DNA was extracted from the caecal contents of these adult mice. The variable region (V1-V2) of the bacterial 16S ribosomal DNA (rDNA) was amplified and metagenotyping using a 454 sequencer was performed. The resulting sequences (3400 reads for each sample) were classified into operational taxonomic units (OTUs) based on sequence similarity (>96% identity). Representative sequences from each OTU were compared with sequences deposited in publicly available 16S and genome databases using BLAST to determine their closest species. As shown in FIG. 14, in +hu mice, OTUs belonging to Bacteroidetes accounted for about 50% of the caecal microbiota community. In contrast, in most OTUs in +huChlo mouse were related to species belonging to Clostridium. In +2x10^4, +2x10^4-re and (+2x10^4)^2 mice, the majority of bacteria consisted of bacteria having 16S rDNA sequence similarities with about 20 species of Clostridia belonging to cluster XvA (also referred to as *C. leptum* group), IV, XVI, and XVIII, listed in FIG. 14.

**EXAMPLE 35**

A meta-analysis of 16S rDNA of caecal contents from mice inoculated with the 23 strains isolated in example 30 (23-mix mice) confirmed the presence of 17 of the 23 strains listed in FIG. 14 and Table 4. To determine whether these 17 strains could induce Treg cells, a mixture of these 17 strains was inoculated into adult GF mice (+17-mix mice). Each bacterial strain was cultured in 2 mL EG liquid media and grown to confluence, and then these cultures were mixed into a 50 mL tube (2 mL per strain = 34 mL). The bacteria were spun down into a pellet and resuspended in 10 mL PBS. A 200 μL aliquot, containing +1x10^8-1x10^7 of each strain, was used to inoculate the adult GF mice. As shown in FIG. 15, when orally administered to adult IQuI, BALB, and B6 mice, the mixture of 17 strains was able to induce Tregs in these three mouse models.

**EXAMPLE 36**

To investigate whether each of the 17 strains defined in example 35 could individually induce Tregs, adult GF mice were monoclonized with one of each of the 17 strains. As shown in FIG. 16, adult GF mice monoclonized with a single strain exhibited low to intermediate levels of Treg. Importantly, no single strain induced Tregs to the same extent as the mix of 17 strains.

**EXAMPLE 37**

To investigate whether subsets of the 17 strains described in example 35 could induce Tregs, randomly selected combinations of 3-5 strains were made: 3-mix, 5 mixA, 5-mix B, and 5-mix C, as shown in table 4, and used to inoculate adult GF mice. As shown in FIG. 17, only the 5-species mixes induced significant increases in the frequency of Treg cells, the magnitude of which was intermediate compared with that observed in +17-mix mice.

**EXAMPLE 38**

To investigate the benefits of administration of the mix of the 17 strains described in example 35 (17-mix), adult SPF mice were orally inoculated with either 17-mix or control media and assessed for the induction of Foxp3+ Treg cells three weeks later. As shown in FIG. 18, there was a significant increase in the frequency of colonic Foxp3+ Treg (CD4) cells after three weeks of treatment.

**EXAMPLE 39**

To evaluate the benefit of administration of 17-mix in an animal model of allergic diarrhea, adult SPF mice were orally inoculated with 17-mix or control media while being treated with ovalbumin (OVA), an inducer of allergic diarrhea. As shown in FIG. 19, the occurrence and severity of diarrhea (diarrhea score) was significantly reduced in mice fed 17-mix relative to control mice.

**EXAMPLE 40**

To evaluate the benefit of administration of 17-mix in an animal model of colitis. Adult SPF mice were orally inoculated with either 17-mix or control media while being treated with trinitrobenzene sulfonic acid (TNBS), a frequently used experimental inducer of colitis. As shown in FIG. 20, SPF 17-mix mice demonstrated lower mortality than control mice on exposure to TNBS.

**EXAMPLE 41**

To evaluate the usefulness of the strains represented in 17-mix as a diagnostic and monitoring tool for ulcerative colitis, we examined the relative abundance of the 17 strains in healthy and ulcerative colitis (UC) human subjects using draft genomic sequences of the 17 strains and publicly available human faecal microbiome genomes generated through the European MetaHIT project. UC subjects (N=20) showed a reduction of the 17 strains compared to healthy subjects (N=15), as shown in FIG. 21.
As has been described above, the compositions and methods described herein make it possible to provide an excellent and well-characterized composition for inducing proliferation or accumulation of regulatory T cells (Treg cells) by utilizing certain human-derived bacteria belonging to the Clostridia class or a physiologically active substance or the like derived from the bacteria. Since the bacterial composition has immunosuppressive effects, the bacterial composition can be used, for example, to prevent or treat autoimmune diseases or allergic diseases, as well as to suppress immunological rejection in organ transplantation or the like. In addition, healthy individuals can easily and routinely ingest the bacterial composition, such as in food or beverage, (e.g., a health food), to improve their immune functions.

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**TABLE 1**

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US 10,342,832 B2
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**TABLE 4**

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**OTU3**

(SEQ ID NO.: 70)

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**OTU9**

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**OTU14**

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**OTU15**

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CATGGGGCTAGAGAAGGCAGCCCA
OTU21

(SEQ ID NO.: 24)

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TGCTTCTGTGTGAAAACCTCCGCTGCTAAGATGACACCGCCTCTGATTAGCTAGT
GAGGGGATACGCCCATCAGATCGGACTGCGGATCGAGGAGATGACGCCA
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OTU23

(SEQ ID NO.: 72)

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CCTTACAGGGGATACAGTTGGCCAATGCTGCTTTAAACCGGACGAGGACG
CCATAGGGTACGCCCATCAGATCGGACTGCGGATCGAGGAGATGACGCCA
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OTU38

(SEQ ID NO.: 73)

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TGCTTCTGTGTGAAAACCTCCGCTGCTAAGATGACACCGCCTCTGATTAGCTAGT
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OTU46

(SEQ ID NO.: 20)

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GCACCGTATGAGGCTAGGATTGATGAGGCTAGCGACACTGCACTGAGGACGCCACCCCA
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CAG

OTU49

(SEQ ID NO.: 74)

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TGCTTCTGTGTGAAAACCTCCGCTGCTAAGATGACACCGCCTCTGATTAGCTAGT
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CATGGGAATAGGAACAGCCCA

OTU55

(SEQ ID NO.: 29)

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OT057
(SEQ ID NO.: 75)
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CATTGGCCTGAGTTGAAATACCTGAAAGTGGTGGCAAGTGATAGTGAAGCTGAGCT
GCCAGGTAAGAGCCCGCTTACAAAGGACCTGAGGAGCCTGAGGAGCCTGAGGAGCCTG
CATTGGGACTGAGAACACAGCCCA

OT073
(SEQ ID NO.: 26)
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CATTGGGACTGAGAACACAGCCCA

OT086
(SEQ ID NO.: 35)
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ACACAGGCAAAGAGGATGAGCTTAAATACACCAGATACACATGGCAAGAGCTGAGCT
CATTGGGACTGAGAACACAGCCCA

OT087
(SEQ ID NO.: 34)
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OT089
(SEQ ID NO.: 76)
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OT092
(SEQ ID NO.: 77)
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OTU101
              (SEQ ID NO.: 78)
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OTU111
              (SEQ ID NO.: 79)
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OTU114
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CATGGCGATGTCGAGAAACATCCCGTGGTGAATGATGATGTCGCTGTTGATAGCGAAGGCTGACGGCCAC
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OTU119
              (SEQ ID NO.: 81)
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ATTGGGACTGAGACACCGCCCA

OTU125
              (SEQ ID NO.: 82)
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OTU131
              (SEQ ID NO.: 83)
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CATGGAAGTGGGGTTGGAATAACATGCGACTGCTGGAAGTTGGCTGCTGATCTGGT
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(SEQ ID NO.: 84)
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(SEQ ID NO.: 85)
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(SEQ ID NO.: 36)
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GCAACACTGCTTCTGACGCTGACGGTACCTTCTGCGTTGAAATAATGGTTGACCTGATGAAACGGCCACAG
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(SEQ ID NO.: 86)
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(SEQ ID NO.: 86)
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ATGCTCCTCGTGTGGTTAAGACATACGGAAGAGGTTCTGGTGTAGTGTG
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OTU182
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CGAGGTAGTAAACCCGAGTATGGAAGATCATCCGTAGCTAGGTTAAGTCTATTATTTAAATGGGAC
GACACGTGACGCGTCGCAACTCGAGACACCCAA

OTU189
(SEQ ID NO.: 91)
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GACACGTGACGCGTCGCAACTCGAGACACCCAA

OTU195
(SEQ ID NO.: 32)
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OTU196
(SEQ ID NO.: 92)
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GACACGTGACGCGTCGCAACTCGAGACACCCAA

OTU199
(SEQ ID NO.: 93)
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GCGAGGGTAAACCCGAGTATGGAAGATCATCCGTAGCTAGGTTAAGTCTATTATTTAAATGGGAC
GACACGTGACGCGTCGCAACTCGAGACACCCAA

GCGGGCAG

OTU202
(SEQ ID NO.: 94)
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GCGAGGGTAAACCCGAGTATGGAAGATCATCCGTAGCTAGGTTAAGTCTATTATTTAAATGGGAC
GACACGTGACGCGTCGCAACTCGAGACACCCAA
CATTGGGACTGAGACACGCCCAG

OTU204

(GEN ID NO.: 96)

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CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA
CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA

OTU211

(GEN ID NO.: 96)

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CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA
CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA

OTU214

(GEN ID NO.: 97)

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CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA

ACATGCGGACTGAGACACGCCCAC

OTU221

(GEN ID NO.: 21)

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CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA
CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA

ACATGCGGACTGAGACACGCCCAC

OTU224

(GEN ID NO.: 99)

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CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA
CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA

ACATGCGGACTGAGACACGCCCAC

OTU225

(GEN ID NO.: 99)

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CTATACAGGGGGATTAAATATGGAGTGGCTGTATGCCGCTCCTATCGCCGCTTTATGCCCTGA
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(SEQ ID NO.: 108)
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CATGGGTCTGCTGAGAAATCCGCCGTTGAAAGACATGTGTGGCTCGGGAG
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GTGCGCGTCTATGCGGCCGCCAGCGAGACTAATCGAGCGCATCGGCTGAGCGCTGAGCG
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(SEQ ID NO.: 109)
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<223> OTHER INFORMATION: OTU337

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gacgacagct ggccgagtgc ctaacacatg caagtcgaac ggacgcttaac ttttgaagtt
   1    2    3    4    5    6
   60

ttcgagatga tgaatgtaag cttcggtgc ggaaggagt gtaaccctgca gcaacgtcgc
   7    8    9   10  11  12  13
   120
tttcagaggg ggatacagc cggaaacgcc tgtaaatacg ggctggtgtg ggccggggac
   14   15  16  17  18
   190
atgcaccctgc aaccaagag gcacagctct gcagacaggg gatcgggtcc attagccagt
   19   20  21
   240
tggtccgggt aaggcgacca aaagcgacga tcggtagccg gacgagagg tggacagcgc
   22   23  24  25  26  27
   300
acattgggac tgagacagcc cccag
   28
   325

<210> SEQ ID NO 31
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU314

<400> SEQUENCE: 31
gatgacacgt ggccgagcgtgc ctaacacatg caagtcgaac ggacgcttaat tttcagaac
   1   2   3   4  5  6  7
   60
ttcgaggaga ggaacaagtg actgacgccc ggacggtgta gtaacgcttg ggcaacgtgc
   8  9 10
   120
tctacagag ggtataaac cttgataatg ctcgatata ccgtctagcc ccacagacgc
   11 12
   180
ctggtgtag tgtagaaaaa tcggcgtgta tgagatgcac gcgcgtctga tttagtagtt
   13 14 15 16
   240
gtggtgggt gcgcgtacca gcgcgtcag tccgctgaggg tggacggcga
   17
   300
acattgggac tggagacagcc cccag
   18
   324

<210> SEQ ID NO 32
<211> LENGTH: 325
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU195

<400> SEQUENCE: 32
gatgacacgt ggccgacggtgc ctaacacatg caagtcgaac ggacgcttatg agatgagasgg
   1   2   3  4   5
   60
ttcgagatggt tgtcgagatgt gatgtgggtc gcgcgtcagc gtaacagcgtg gtaacagcgc
   6  7
   120
tctacactgg ggcaacgaca cttgatataa ctgcgataac gcacagtacc
   8  9 10
   190
goatgtacgc gtgaatgaaa tccggtgtgt gccgatgacg tccgctagt tttagccagt
   11 12
   240
tggtccggtta cagcccaacgc aagcgagcg tcaagacgac gctgagaggg tggacgcggc
   13
   300
acattgggac tgagacacgac cccaa
   14
   325

<210> SEQ ID NO 33
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU306
<400> SEQUENCE: 33

gatgaacgct gcggcgctgc ttaacacatg caagtcgaac gaaagcgaatt aacggaagtt 60
ttcggatgg aaatgatagtacaggtggc gaaagcgtga tcgaagcgtg ggttaacgc 120
cattgaatgg gcggcgaacactacattagtaaacctaggtagcaacactagttccagt 180
gatgcatac gccgatagtaaaggtcgctgacacagcgtaagcggtgtagtaaagttattaggtg 240
tgtagtacgta acgcggctacccagcgcggtggacagtggggtgtgtagtggtgtgcc 300
aacggtggc ttagacacccg cccca 324

<210> SEQ ID NO 34
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTUB7

<400> SEQUENCE: 34

gatgaacgct gcggcgctgc ttaacacatg caagtcgaac gaaagcgaatt aacggaagtt 60
ttcggatgg aaatgatagtacaggtggc gaaagcgtga tcgaagcgtg ggttaacgc 120
cattgaatgg gcggcgaacactacattagtaaacctaggtagcaacactagttccagt 180
gatgcatac gccgatagtaaaggtcgctgacacagcgtaagcggtgtagtaaagttattaggtg 240
tgtagtacgta acgcggctacccagcgcggtggacagtggggtgtgtagtggtgtgcc 300
aacggtggc ttagacacccg cccca 324

<210> SEQ ID NO 35
<211> LENGTH: 325
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTUB6
<220> FEATURE:
<221> NAME/KEY: mrc_feature
<222> LOCATION: (49) (49)
<223> OTHER INFORMATION: n is a, c, g, or t

<400> SEQUENCE: 35

gatgaacgct gcggcgctgc ttaacacatg caagtcgaac gaaagcgaatt tgaagcaag 60
ttcggatgg aaatgatagtacaggtggc gaaagcgtga tcgaagcgtg ggttaacgc 120
acccatatgc gcggcgaacactacattagtaaacctaggtagcaacactagttccagt 180
gcgatgtcct gcggcgaacactacattagtaaacctaggtagcaacactagttccagt 240
tgtagtacgta acgcggctacccagcgcggtggacagtggggtgtgtagtggtgtgcc 300
aacggtggc ttagacacccg cccca 325

<210> SEQ ID NO 36
<211> LENGTH: 323
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU152

<400> SEQUENCE: 36

gatgaacgct gcggcgctgc ttaacacatg caagtcgaac gaaagcgaatt aaggaagtt 60
ttcggatgg aaatgatagtacaggtggc gaaagcgtga tcgaagcgtg ggttaacgc 120
cgtagtggtgaactactacattagtaaacctaggtagcaacactagttccagt 180
atgggtcttg tgtgaaact cccgtggtac agagttgcc cggctctgat tagccagttg
324

goaggttaac ggccttacaa agcagcgcgc actgacccggct ctggacagggt gaacgggccac
300
attggacagt agacacggcc caa
323

<210> SEQ ID NO 37
<211> LENGTH: 326
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU253

<210> SEQ ID NO 38
<211> LENGTH: 326
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU259

<210> SEQ ID NO 39
<211> LENGTH: 127
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU281

<210> SEQ ID NO 40
<211> LENGTH: 222
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU288
<400> SEQUENCE: 40

gatgaacgtg gcgccgcgtgc ctaacacatgc caacgctgaac ggagttatgc agaggaagtt 60
tccgatgga atcgctaaat ctagcgcggt gcgggtgtcg ggacgcctgc gcc 120
cggattcgg gcggtattgc tcgagaatt gcacgaatgtg gcgcctgtgc gcaacgtgcagc 180
atgagaggtc gttgaaatgg tggcggtgc ggaggtcttc gctggtctgtgtgc ggtgcagct 240
goagagtcgg gcgttcagct ggtgcggcgtgc atggggtgtc cttgagggg gcggcagc 300
atcggctag agacacgccca 322

<210> SEQ ID NO 41
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU334

<400> SEQUENCE: 41

gatgaacgtg gcgccgcgtgc ctaacacatgc caacgctgaac ggagttatgc ataggaagtt 60
tccgatgga atcgctaaat ctagcgcggt gcgggtgtcg ggacgcctgc gcc 120
cggattcgg gcggtattgc tcgagaatt gcacgaatgtg gcgcctgtgc gcaacgtgcagc 180
atgagaggtc gttgaaatgg tggcggtgc ggaggtcttc gctggtctgtgtgc ggtgcagct 240
goagagtcgg gcgttcagct ggtgcggcgtgc atggggtgtc cttgagggg gcggcagc 300
atcggctag agacacgccca 324

<210> SEQ ID NO 42
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU359

<400> SEQUENCE: 42

gatgaacgtg gcgccgcgtgc ctaacacatgc caacgctgaac ggagttatgc agaggaagtt 60
tccgatgga atcgctaaat ctagcgcggt gcgggtgtcg ggacgcctgc gcc 120
cggattcgg gcggtattgc tcgagaatt gcacgaatgtg gcgcctgtgc gcaacgtgcagc 180
atgagaggtc gttgaaatgg tggcggtgc ggaggtcttc gctggtctgtgtgc ggtgcagct 240
goagagtcgg gcgttcagct ggtgcggcgtgc atggggtgtc cttgagggg gcggcagc 300
acattgggggc tggagaacgg ccgc 324

<210> SEQ ID NO 43
<211> LENGTH: 321
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU362

<400> SEQUENCE: 43

gatgaacgtg gcgccgcgtgc ctaacacatgc caacgctgaac ggagttatgc cgatgaagtt 60
tccgatgga tttaatacgc acctagcgcc gccggtttgc gcgctgcggcg gtaacgctgc gcaacgtgcagc 120
cggattcgg gcggtattgc tcgagaatt gcacgaatgtg gcgcctgtgc gcaacgtgcagc 180
atgagaggtc gttgaaatgg tggcggtgc ggaggtcttc gctggtctgtgtgc ggtgcagct 240
goagagtcgg gcgttcagct ggtgcggcgtgc atggggtgtc cttgagggg gcggcagc 300
}
tgagactgag acacgccca a

<210> SEQ ID NO 44
<211> LENGTH: 323
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU367

<400> SEQUENCE: 44

gatgaacgc ggcggcgtgc ctaacacatg caagtgaac gaaagtacac agaggaagtt  60
tcggatgga atcgttataa cttatggtcgc gacggtgtag taacgcgtgg gaacacgccc 120
cgtacgccc ggtataacat tagaaatagg tgtaataacc gcataaggc acagctcacc 180
tagaaacagt gtgaataacct cggtaggtac agaatggtcc cgcgctgat tagcgaattg 240
gcaggctaac ggcctaccaaa agcgaagcgg ctagcggg gcctgaggt gcaacgccac 300
atgggactg agacacgccac cag  323

<210> SEQ ID NO 45
<211> LENGTH: 17
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, total bacteria

<400> SEQUENCE: 45

ggtgaatacg ttcgccg  17

<210> SEQ ID NO 46
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, total bacteria

<400> SEQUENCE: 46

tacggttacc tgtaaagac tt  22

<210> SEQ ID NO 47
<211> LENGTH: 20
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, Clostridium cluster XIVa (Clostridium coccoides subgroup)

<400> SEQUENCE: 47

aaatgacggt acctgactaa  20

<210> SEQ ID NO 48
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, Clostridium cluster XIVa (Clostridium coccoides subgroup)

<400> SEQUENCE: 48

cottgatctc ccctttgagc aa  22

<210> SEQ ID NO 49
<211> LENGTH: 16
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Primer, Clostridium cluster IV (Clostridium leptum)

SEQUENCE: 49
cttcggcgct gcaga tta

SEQ ID NO 50
LENGTH: 26
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Primer, Clostridium cluster IV (Clostridium leptum)

SEQUENCE: 50
gat taa acc acata c tca ctgct t

SEQ ID NO 51
LENGTH: 18
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Primer, Bacteroides

SEQUENCE: 51
gagag gag tcoc ccac

SEQ ID NO 52
LENGTH: 19
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Primer, Bacteroides

SEQUENCE: 52
cgtcttgg ctg ttcag

SEQ ID NO 53
LENGTH: 20
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Primer, Bifidobacterium

SEQUENCE: 53
cgggtgaga tctgctgacc

SEQ ID NO 54
LENGTH: 18
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Primer, Bifidobacterium

SEQUENCE: 54
tga tag acgc caga coca

SEQ ID NO 55
LENGTH: 50
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Modified primer 6F
FEATURE:
NAME/KEY: misc_feature
LOCATION: (30) .. (31)
OTHER INFORMATION: barcode

SEQUENCE: 55
ccatctcctc ctgcgtgtgct tccgactcag agrgtttggt ymtggctcag

SEQ ID NO: 56
LENGTH: 49
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Modified Primer 338R

SEQUENCE: 56
cctatccct cgtgccttg gcagtctcag tgcgtgcctc cgtaggagt

SEQ ID NO: 57
LENGTH: 19
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: KOD FX (TOYOBO), 16S rRNA gene-specific primer, 8P

SEQUENCE: 57
gagtttgcct tmtgctcag

SEQ ID NO: 58
LENGTH: 16
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: KOD FX (TOYOBO), 16S rRNA gene-specific primer, 519R

SEQUENCE: 58
attacccgg cgctcg

SEQ ID NO: 59
LENGTH: 21
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: KOD FX (TOYOBO), 16S rRNA gene-specific primer, 513R

SEQUENCE: 59
acggctacct tgttacgct t

SEQ ID NO: 60
LENGTH: 20
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: 16S rRNA gene-specific primer

SEQUENCE: 60
agagtttggat cmctggtcag

SEQ ID NO: 61
LENGTH: 16
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: 16S rRNA gene-specific
attacgccg ckgctg

<210> SEQ ID NO 61
<211> LENGTH: 16
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, total bacteria

<400> SEQUENCE: 61

<210> SEQ ID NO 62
<211> LENGTH: 20
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, total bacteria

<400> SEQUENCE: 62
ggtaatacg ttcccgg

<210> SEQ ID NO 63
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, total bacteria

<400> SEQUENCE: 63
tacggtacct tgttacgat tt

<210> SEQ ID NO 64
<211> LENGTH: 20
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, Clostridium cluster XIVa (Clostridium coccoidees subgroup)

<400> SEQUENCE: 64

aasngctgg acttgcttas

<210> SEQ ID NO 65
<211> LENGTH: 22
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Clostridium cluster XIVa (Clostridium coccoidees subgroup)

<400> SEQUENCE: 65

ctttgagttt cattcttcgg ac

<210> SEQ ID NO 66
<211> LENGTH: 16
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, Clostridium cluster IV (Clostridium leptum)

<400> SEQUENCE: 66
gcacacgcag tggagt

<210> SEQ ID NO 67
<211> LENGTH: 16
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, Bacteroides

<400> SEQUENCE: 67
gagagagagag tccccccac
<210> SEQ ID NO 68
<211> LENGTH: 19
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, Bacteroides

<400> SEQUENCE: 68
gctacttgg ctgttccag 19

<210> SEQ ID NO 69
<211> LENGTH: 18
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Primer, Clostridium cluster IV (Clostridium leptum)

<400> SEQUENCE: 69
cctcctcgt ttgttcaaa 18

<210> SEQ ID NO 70
<211> LENGTH: 323
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU3

<400> SEQUENCE: 70
gatgacgct gccgcggctgc ttaacacatg caagtccagc gaagcactaa gcggatttc 60
ttcggatgga agcttcttctg actgagccgc ggacggtgag taacgtcttg gtaactgtcgc 120
atcatacgct ggataacagt tagaaatagc tgtgtaaacc gctataagcg acaggacgacg 180
atggtctggt gtagaaaact cccgtggtat gagatggacc cgctgctgtat tagttgatg 240
gagggtaacgc gocacacgac gccgacgttc agtagccggc ctgaggggtt gacgccgacc 300
atggtgactg agacacggtcc cag 323

<210> SEQ ID NO 71
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU15

<400> SEQUENCE: 71
gatgacgct gccgcggctgc ttaacacatg caagtccagc gaagcactaa gcggattttc 60
ttcggatgga agcttcttctg actgagccgc ggacggtgag taacgtcttg gtaactgtcgc 120
tctatacgct ggataacagt tagaaatagc tgtgtaaacc gctataagcg acaggacgacg 180
catggtctggt gtagaaaact cccgtggtat gagatggacc cgctgctgtat tagttgatg 240
gagggtaacgc gocacacgac gccgacgttc agtagccggc ctgaggggtt gacgccgacc 300
catggtgactg agacacggtcc cca 324

<210> SEQ ID NO 72
<211> LENGTH: 224
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU23
gatgaacgct ggcggcggtgc ttaacacatg caaagtcgac gaagcatattt ggaaggaagt 60
ttccggatgg aatcccttaa tgaatgcagt ggcggcggtg gtaaactcg gtaaggaact 120
cctcataca ggggtagaag acgtggaac acagctaatg acgcctaaag ggcacagaa 180
cagatgttc ttgggtgaag atctggcccgt ataggggattt cccctgtctg gtagtctgt 240
tggcgggttaa aggcctcaac gaagcgcact cagagtgcgg cttgagagag tggacggcaca 300
cattgggact gagaacaggg ccac 324

<210> SEQ ID NO 73
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU38

<400> SEQUENCE: 73

gatgaacgct ggcggcggtgc ttaacacatg caaagtgcag gcagcgcgattt caaataggtt 60
ttccggatgg aatcccttaa tgaatgcagt ggcggggtg gtaaactcg gtaaggaact 120
cctcataca ggggtagaag acgtggaac acagctaatg acgcacaaag ggcacagaa 180
cagatgttc ttgggtgaag atctggcccgt ataggggattt cccctgtctg gtagtctgt 240
ggcggggtaa aggcctcaac gaagcgcact cagagtgcgg cttgagagag tggacggcaca 300
cattgggact gagaacaggg ccac 324

<210> SEQ ID NO 74
<211> LENGTH: 329
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU49

<400> SEQUENCE: 74

gatgaacgct ggcggcggtgc ttaacacagc caaagcgcac gaagccatattt aatgacagt 60
ttccggatgg tttttgtgttg actagtttgc gcacgggtgga tgaacgcgtg gtaaactcg 120
cctcatacg ctcctaccaag actgtaatc attccatagc gcacactttc gacacgtaa 180
goatgtagc ttgaaatatg ccgccctgtg tgaagtggag tccgctttc tttcgagatg 240
tggccggggtaa aacgcgcctac caagcgcact cagagtgcgc atctggaggag tggacggcaca 300
cacatattggg aatgagacac gcgccccc 328

<210> SEQ ID NO 75
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU57

<400> SEQUENCE: 75

gatgaacgct ggcggcggtgc ttaacacatg caaagtgcag gcagcgcgattt cgatgaagtt 60
ttccggatgg tttgaatcgc actagcgcc gcacgggtgga tgaacgcgtg gtaaactcg 120
cctcatacg ctcctaccaag actgtaatc attccatagc gcacactttc gacacgtaa 180
goatgtagc ttgaaatatg ccgccctgtg tgaagtggag tccgctttc tttcgagatg 240
ggcggggtaa aacgcgcctac gaagcgcact cagagtgcgc atctggaggag tggacggcaca 300
cattgggact gagacacggc cca

<210> SEQ ID NO 76
<211> LENGTH: 327
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU89

<400> SEQUENCE: 76

gatgaaacgt ggcggcgtgc ttaacacagt caagtcgaaac gaaacctttt ggaaggaaatg
60
ttcggatgta atcgggtata acttagtggc gcggcgggta gtaacgcgtg ggaacactgc
120
cctgtacgcgg ggacacacac ttaagatag tgaatcaaac cgcaataacgc acggaacacg
180
catggctcccg tggcagaaaa ttcggcggtgt cacgagtccg tccccggtc tcctagccagtt
240
tgccccgggt aacagcctca ccaagacacag gatactgtcgc cggcgtgaga ggtgtaaaccg
300
cocactctgg actgacacag gcggccca
327

<210> SEQ ID NO 77
<211> LENGTH: 322
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU92

<400> SEQUENCE: 77

gatgaaacgt ggcggcgtgc ctaacacagt caagtcgaaac ggaattactgc agagggactt
60
ttcggatgta atcgggtata acttagtggc gcggcgggta gtaacgcgtg ggaacactgc
120
cctgtacgcgg ggacacacac ttaagatag tgaatcaaac cgcaataacgc acggaacacg
180
catggctcccg tggcagaaaa ttcggcggtgt cacgagtccg tccccggtc tcctagccagtt
240
tgccccgggt aacagcctca ccaagacacag gatactgtcgc cggcgtgaga ggtgtaaaccg
300
cocactctgg actgacacag gcggccca
322

<210> SEQ ID NO 78
<211> LENGTH: 323
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU101

<400> SEQUENCE: 78

gatgaaacgt ggcggcgtgc ctaacacagt caagtcgaaac gaaaccttta ataggaagtt
60
ttcggatgta atcgggtata acttagtggc gcggcgggta gtaacgcgtg gataacactgc
120
cctgtacgcgg ggacacacac ttaagatag tgaatcaaac cgcaataacgc acggaacacg
180
catggctcccg tggcagaaaa ttcggcggtgt cacgagtccg tccccggtc tcctagccagtt
240
gcgcgggttta cgcacacacg acgagcagat gactgacgcc cctgagaggg tcaccggcaca
300
cactggact gagacacggc cca
323

<210> SEQ ID NO 79
<211> LENGTH: 322
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU111

<400> SEQUENCE: 79

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cggattggcac tagacacaggg cccaa

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<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU131
<220> FEATURE:
<223> OTHER INFORMATION: n is a, c, g, or t
<400> SEQUENCE: 83

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tcaacactgg gggttaacag ttagaataatgc tggtcaataac cggataagcc cacagtgccg 180
catggcgtgc tggtaaasac tccggtgtgtg tgtagatgatc cccggtctga ttaggacgat 240
ggcggtaacgg gcacacgacag gcagcagatca tgacgagggc tgaacggagggt nacoggcaca 300
ttggacgta gacacggccc aa 322

<210> SEQ ID NO 84
<211> LENGTH: 325
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU137
<400> SEQUENCE: 94

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tccggatgga atcttgatggt actgaagtggc ggacggttga acctgcgtg gataacgcctg 120
tcaacactgg gggttaacag ttagaataatgc tggtcaataac cggataagcc cacagtgccg 180
ggcggtaacgg gcacacgacag gcagcagatca tgacgagggc tgaacggagggt nacoggcaca 240
ttggacgta gacacggccc aa 325

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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU144
<400> SEQUENCE: 85

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tttgcgttga cagcggtacc acgcagcgg ggaacggttga gtaacgcgtg ggtagacgctc 120
tcaacactgg gggttaacag ttagaataatgc tggtcaataac cggataagcc cacagtgccg 180
catggcgtgc tggtaaasac tccggtgtgtg ttgtagatgatc cccggtctga ttaggacgat 240
ggcggtaacgg gcacacgacag gcagcagatca tgaagggcgt ggacgagggc tgaacggagggt 300
ttggacgta gacacggccc aa 325

<210> SEQ ID NO 86
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU161
<400> SEQUENCE: 86

<400> SEQUENCE: 87

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cctacactcg gcggacaacg ttagaaatga ctgtctaatacc gctataaacc cacagtacgc 180
catgtgaac tgtggaaaaac tcctggggtca cagatggtgc cggctctgtga ttcaacccag 240
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cattggagact gcagaacgagc ccaa 324

<400> SEQUENCE: 88

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cctacactcg gcggacaacg ttagaaatga ctgtctaatacc gctataaacc cacagtacgc 180

<400> SEQUENCE: 89

<400> SEQUENCE: 90

<400> SEQUENCE: 91

<400> SEQUENCE: 92

<400> SEQUENCE: 93
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<210> SEQ ID NO: 90
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU182

<400> SEQUENCE: 327

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ggggaacgggg tgaagtaac ataagtaacc tgccccttttc agggggataag ccattggaa 120
cagagctaa gacccgctag gtaaaagtaac gccaatggttaa gttttataaa agtgcccaag 180
caccagggagtagcttgat atgctgagtctagtgcttgt aggttaacggg tcacaagggc 240
gacgtgctg agccgacactg agagggtgac ccgcaacact gcggactgaga caagggc 300

<210> SEQ ID NO: 91
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU189

<400> SEQUENCE: 91

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tcactcgg ggcacaagct tagaatgac tgctaatacc gecataagcgc acagcctcac 180
atgcagagt gtaaaaaact gcccgggtct agataagcct ccggtctgtg tagcaaatgtg 240
gcagggtac gcctaacac agctgagctg atctagcgcgc ctgagaggtt gaagggccac 300
attggaacct agacaagggcc cag 323

<210> SEQ ID NO: 92
<211> LENGTH: 322
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU196

<400> SEQUENCE: 92

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cattcaggg gacaaaaagct tggcaacagct ggtaataacc gcgtagacaca ctagtggggc 180
atcaggcagc tgcccaatag gtagactgca aagatggcgtc cgctgttccc agtagctgtg 240
gcgggggtac gcctaacaca ggcagcgtag cggtagcggc ctgagaggtt gacgagggcc 300
attggaacct aggataagggcc ca 322

<210> SEQ ID NO: 93
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU199

<400> SEQUENCE: 93

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cgaagcata gaccgcatag gtaacagcag tgggtggtga cctgatattaa aagtagctcctca 180
aagaacactag tgaagatcag actcatggtc cattagcagg tgggctgggt aacgccccac 240
caacgcaac gatacctcag cgaacttcgaga ggtgacccgg ccacactggg actgagacac 300
ggccccag  307

<210> SEQ ID NO 94
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU202

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ttcgatggga aagtgaatttg acgacttgcc ggacgggtga tgaacgcttg gttaacctgc 120
cctgctttcg ggaacaaag tctagaaatc ctgctaatc ccgataagcg caacagatcc 180
cattgatacg tgtaaaaacct ccgctttgga caaagttgcc cgcgtctgga ttagctggtt 240
ggacggttcc ggcccccaaca agggcagagc cagtagcgg gctgagagg ggaaacggcca 300
cattgaggct gacacacggc cca 324

<210> SEQ ID NO 95
<211> LENGTH: 323
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU204

<400> SEQUENCE: 95
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ttcgatggga aagtctcggtt acgacttgcc ggacgggtga tgaacgcttg gttaacctgc 120
cctcataaggggtaacatg ttagaaatc ccgataagcg caacagatcc 180
cattgatacg tggaaaaaccc cccggtttgga cccgctttgga ttagctggtt 240
ggtaaagttcc ccgctttgcac agggcaagac cagtagcgg gctgagagg ggaaacggcca 300
cattgaggct gacacacggc cca 323

<210> SEQ ID NO 96
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU211

<400> SEQUENCE: 96
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ttcgatggga ttagaaatcag acctagcggc ggacggtgta tgaacgcttg gttaacctgc 120
cctcataaggggtaacatg ttagaaatc ccgataagcg caacagatcc 180
cattgatacg tggaaaaaccc cccggtttgga cccgctttgga ttagctggtt 240
ggacggttcc ggcccccaaca agggcagagc cagtagcgg gctgagagg ggaaacggcca 300
cattgaggct gacacacggc cca 324

<210> SEQ ID NO 97
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE: 
<223> OTHER INFORMATION: OTU214

<400> SEQUENCE: 97

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ttccggttga acctttaggta actgagtggc gagacgggtga tgaacgctgt ggtaacacgtc 120
cctacaagg ggagtaaaca gtttaaattg actgtaataa ccgctatagc gcacagggct 180
gcatggccttg gtgtaaaas ctcgcttggt atgagatgga ccgctgtctg attagcctagt 240
tggaggggta acggccccaco aagggcagta tcagtaacgc gctgtaagagg gtgaacggcct 300
acattgggac tgacacagcg ccca 324

<210> SEQ ID NO: 98
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<213> ORGANISM: Artificial Sequence
<220> FEATURE: 
<223> OTHER INFORMATION: OTU224

<400> SEQUENCE: 98

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cctacaagg ggagtaaaca ttagaataag gtctaatata ccgctatagc gcacacgtttca 180
catgaacag tggtaaataac tccgggctgta cagacgtgtc cccgtcctgta ttacccagt 240
gacggggcta cgccctaccg aacgcacagct tcagtaacgc gctgtacggg tgaacggcct 300
cattgggac gagaacacgg cca 323

<210> SEQ ID NO: 99
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE: 
<223> OTHER INFORMATION: OTU225

<400> SEQUENCE: 99

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ccttcgaagg ggagcctaac acttactata gttctaatata ccgctatagc gcacacgcttc 180
acattaggca gtgtgaaataa ctcgccgtgtgt acaggatgtc ccgctgtctg attagcctagt 240
tggcagggta acggccctacc aacgcagacga tcagtaacgc gctgtacggg tgaacggcct 300
acattgggac tgacacagcg ccca 324

<210> SEQ ID NO: 100
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<213> ORGANISM: Artificial Sequence
<220> FEATURE: 
<223> OTHER INFORMATION: OTU227

<400> SEQUENCE: 100

 gagtaacgct ggccggctgct ctaacacatg caagtcaaac gsaacaattg aaggaagttt 60
tggaggtggc ttcggttagc tgaagttggg acgggagtgt aecgccgtga tatacctgtc 120
cacccgggg gataacagt tggaaaaact gtaaatcagc ctaacgcgca ccgctgcgca 180
tgtagcagctg taaacaactt cgggtggtgtg agatggactt gcgcctgttt gacgcgtttg 240
cggggtaacg gcocacaaac gcagcagcatc gtacggccct tgagaggtgtg acgggcacac 300
ttgggacgta gcacagggcct ca 322

<210> SEQ ID NO 101
<211> LENGTH: 323
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU246

<400> SEQUENCE: 101
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cttgaggtga atcggggttga cttatggtgc gcacgggtgag taacgctgtg ccaaaccctgc 120
cataccaggg ggataacgcg tggaaacggg tggtaattac cctaagacgc acagaatgcg 180
agatcgttgc ttgaaaagct cccggcagttg agatgctgctgcc gcgggtggtc tagatgtgctg 240
gcgggctac ggccacacaa gcggcagagt atagccggcc ttgacagaagt gcagggccac 300
atgaggaacg tagacacacg ccaa 323

<210> SEQ ID NO 102
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU262

<400> SEQUENCE: 102
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tttgaggtga tttgaggtag atcgagtgcg gacgggtgta gcatacgttg ctaacagtgc 120
cataccatgg ggacacacacct ttgaaatattc cttgataatac cggcataagg cacaggtacgc 180
catggtgtgc tggaaacacc tctggtggtcg tggagattag gcgggtgcttg ttgacacgat 240
gcggggtgtta gcggccacaa acagcagigcg caggtagcct cgctgaggg tgtcggcgccaa 300
cattggggac tttgacacg gcacca 325

<210> SEQ ID NO 103
<211> LENGTH: 326
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU268

<400> SEQUENCE: 103
gatgaaagct ggccgcgtgct ctaacacatg caagtcgaag gcagcgctgt tttcgaatcc 60
tttggaggtga gacacgctggt actcggggcg gacgggtgga gcacggctgtgc gcgaactgcg 120
catacaggg ggataaacag ttgaaatagta cttgataatac cgcataagcg cacaggccag 180
catggggtgag tgtaaaacac cggcgtggtcttgtg tttgtggtatgc tttggtggtgag 240
gcgggctgga aacgctgcaaac gcggcagagt acagtccgaa cttgagaagg gttggccgaac 300
cattggggac tttgacacg ccacca 326

<210> SEQ ID NO 104
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU269
<400> SEQUENCE: 104

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ttcctgaatg atctgagtgc ggaagggcgc gagaagtgcg ctaacagctg gataacctgc  
120
tcaacatgc ggagacgaca gttgaaataa gactctaat aggcctaaag gcacacagct  
180
cgctaagctg gattgtaaaa actaacggtg gtgtgagatg gttcagcgct gattgctca  
240

gttgcgcggg taaacgccca ccaaaacgacgc gttcagctgc gcacccgaa gggtgccgag  
300
coscacgttg gcatcagacgccccc  
329

<210> SEQ ID NO 105
<211> LENGTH: 126
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE: 
<223> OTHER INFORMATION: OTU277

<400> SEQUENCE: 105

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ttcctgaatg atctgagtgc ggaagggcgc gagaagtgcg ctaacagctg gataacctgc  
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tcaacatgc ggagacgaca gttgaaataa gactctaat aggcctaaag gcacacagct  
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cgctaagctg gattgtaaaa actaacggtg gtgtgagatg gttcagcgct gattgctca  
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tggagggca aggggcaccc aagggcagca ttgatagcgg gctgtgagag ggtgaagcgg  
300
acgattgga ctagacaagc gcgccag  
326

<210> SEQ ID NO 106
<211> LENGTH: 323
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE: 
<223> OTHER INFORMATION: OTU279

<400> SEQUENCE: 106

gatgacagct ggccgcatg ctaacacatg caagtcgaac gaagtgagac agaggaaggt  
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ttcctgaatg atctgagtgc ggaagggcgc gagaagtgcg ctaacagctg gataacctgc  
120
tgcacctgag ggataaactg ttgatagcgg gctgtgagag ggtgaagcgg  
180
caggtgtgtg gttgtaaaa actaacggtg gtgtgagatg gttcagcgct gattgctca  
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gcatcagctgc gcacacagcctgactgagag gctgtgagag ggtgaagcgg  
300
catgggact gacacagcgc cca  
323

<210> SEQ ID NO 107
<211> LENGTH: 315
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE: 
<223> OTHER INFORMATION: OTU280

<400> SEQUENCE: 107

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ttttacagct gatgccccag ggggtagtga atccataagc aacctgccca tgggccgctgg  
120
taatgtagatg gaagctgagt ttaaaccgct ctattctgtg aaggtctaat  
180
taagctgagct gctttgtgaaa agagacgctg gctttgtgaaa agagacgctg  
240
gcgcaccc agggctgcag cttcagccgg gcacagccgc gttgagcggc  
300
tagacacacgc ccaaa  
315
<210> SEQ ID NO 108
<211> LENGTH: 325
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU286

<400> SEQUENCE: 109

<210> SEQ ID NO 109
<211> LENGTH: 329
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU287

<400> SEQUENCE: 109

<210> SEQ ID NO 110
<211> LENGTH: 328
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU297

<400> SEQUENCE: 110

<210> SEQ ID NO 111
<211> LENGTH: 256
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU303

<400> SEQUENCE: 111

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**<212> TYPE: DNA**

**<213> ORGANISM: Artificial Sequence**

**<220> FEATURE:**

**<223> OTHER INFORMATION: OTU0304**

**<400> SEQUENCE: 112**

| ATGAACTGT GGGCGGAGGC CTCAACGACT CGAAGTCGAGA AGAACGTTGCG | 60 |
| TGGCTTCTG TAGAGGCTGG GGCGGCTGAG TTATGTCTG GAACCTGCCC GATGGAGGGG | 120 |
| GATAACACTG GGAAGCGTGAA GTCAACCTG CAGACACTG CAGACACCA AAGGAGGACC | 180 |
| TTGAGCTTGG TCTGCACTG CGAATGGTCG ATGGATATTG CTAGCTGG GTAGAAAAG | 240 |
| GGCCTACTG GAGAGCCAGC CAGTACGGA CACAAGACGC GACGCGCAGA | 300 |
| CAGCGCTGAC | 326 |

**<210> SEQ ID NO 113**

**<211> LENGTH: 325**

**<212> TYPE: DNA**

**<213> ORGANISM: Artificial Sequence**

**<220> FEATURE:**

**<223> OTHER INFORMATION: OTU0307**

**<400> SEQUENCE: 113**

| GATGAACTGT GGGCGGAGGC CTCAACGACT CGAAGTCGAGA AGAACGTTGTT | 60 |
| TTCGGATTTG ATATGAGGAA TGAACACTG GATAACCTGCG GAGAGCCAG | 120 |
| CCTCACTCTG GCCGACACTG TAGAAGATG CTCTGACACT GAGAGCCAG | 180 |
| GCTGGTAGG CTGAGAAAG CGCAAGAGT GCTGCGCTTGG ATTAGCCAGT | 240 |
| TTGGGATTAG GGCAGGACGC CAGTACGGA CACAAGACGC GACGCGCAGA | 300 |
| GATGAACTGT GAGAGGACC GCCGAG | 325 |

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**<211> LENGTH: 123**

**<212> TYPE: DNA**

**<213> ORGANISM: Artificial Sequence**

**<220> FEATURE:**

**<223> OTHER INFORMATION: OTU0312**

**<400> SEQUENCE: 114**

| GATGAACTGT GGGCGGAGGC CTCAACGACT CGAAGTCGAGA AGAACGTTGTT | 60 |
| TTCGGATTTG ATATGAGGAA TGAACACTG GATAACCTGCG GAGAGCCAG | 120 |
| CTGGGACGG CATAACTG TGAAGATG CTCTGACACT GAGAGCCAG | 180 |
| GATGAACTGT GATGAACTGT GATGAACTGT GATGAACTGT GATGAACTGT | 240 |
| GGCCTACTG GAGAGCCAGC CAGTACGGA CACAAGACGC GACGCGCAGA | 300 |
| ATGGGGAAT GAGAGCCAG GCCGAG | 323 |

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**<211> LENGTH: 301**
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<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU313

<400> SEQUENCE: 115

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ggcgaaccgg tgaagataac ataaagttac ggctctttaa aggggatata cattggaaa 120
cgatgctaa gacgcatag tgaagatata cgcatggttaa gtatttataa gtcgcaaggg 180
actgctagag gatgagacta tgcgccattg ctaaagttgt cagagtaaag ctcaacacaggg 240
cgacgagtcg tagccgacct gagaggtgag cccgccaacac tgtgacagtg aacagggcca 300
a 301

<210> SEQ ID NO 116
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
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<223> OTHER INFORMATION: OTU319

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cgtacccgg gatgcataac tagaaatgac tggataacag gcataaggoct aagagctgac 180
atggtgctgt gtaaacacag cccgggattc gatagttgac cgcgtttcag tagccgacctg 240
gccgggtaac gcggccaaca agccgctatc aagtcgcgcgt ctcgagaggt gcggccaca 300
tggggatga gacagggcco aa 322

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<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU326

<400> SEQUENCE: 117

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cgatgaagcc tggtaaaata cccgggtttg gagagtattt ggcgggacag cgcgtgagtg ttagccgactt 240
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<210> SEQ ID NO 118
<211> LENGTH: 324
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: OTU329

<400> SEQUENCE: 118

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tccgggatgc gaacatcag gacctaaagct cgtcattgta acgagctgct gagagttcgg 180
What is claimed is:
1. A pharmaceutical composition comprising a purified bacterial mixture consisting of bacteria comprising 16S rDNA sequences of at least 95% homology to SEQ ID NO:19, SEQ ID NO:20, SEQ ID NO:21, SEQ ID NO:22, SEQ ID NO:24, SEQ ID NO:25, SEQ ID NO:27, SEQ ID NO:30, SEQ ID NO:31, SEQ ID NO:32, SEQ ID NO:33, SEQ ID NO:34, SEQ ID NO:39, SEQ ID NO:40, SEQ ID NO:41, and SEQ ID NO:42, wherein the bacterial mixture induces the proliferation and/or accumulation of regulatory T cells, and one or more enteric polymers.
2. The pharmaceutical composition of claim 1, wherein the bacteria are human-derived bacteria.

3. The pharmaceutical composition of claim 1, wherein the bacteria are isolated from a chloroform-treated fecal sample.
4. The pharmaceutical composition of claim 1, wherein the bacteria are isolated from a heat-treated fecal sample.
5. The pharmaceutical composition of claim 1, wherein at least a portion of the bacteria are in spore-form.
6. The pharmaceutical composition of claim 1, further comprising a pharmaceutically acceptable excipient.
7. The pharmaceutical composition of claim 1, wherein the pharmaceutical composition is formulated for oral administration.